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(57) Abstract

Disclosed is a polynucleotide comprising a chimeric viral RNA which contains: a 5' nontranslated region (5' NTR), an open reading frame (ORF) region, and a 3' nontranslated region (3' NTR) wherein at least one of said regions is chimeric. The chimeric region comprises a first nucleotide sequence from a pestivirus in operable linkage with a heterologous nucleotide sequence. The chimeric viral RNA is replication—competent. Preferably the pestivirus sequence is from a bovine viral diarrhea virus and the heterologous nucleotide sequence is from a hepatitis C virus. Also disclosed are a method for identifying compounds having antiviral activity agaisnt hepatitis C virus, a genetically—engineered chimeric RNA virus and a vaccine against bovine viral diarrhea virus.

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Chimeras of Hepatitis C Virus and Bovine Viral Diarrhea Virus

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Related Applications

This application claims priority to, and incorporates herein in its entirety, U.S. 60/082,964 filed April 24, 1998.

10 Background of the Invention

(1) Field of the Invention

This invention relates generally to the development of therapies for treating hepatitis C virus (HCV) and bovine viral diarrhea virus (BVDV) and more particularly to the identification of such therapies using chimeric viruses comprising a genomic sequence derived from HCV and bovine viral diarrhea virus (BVDV).

(2) Description of the Related Art

The Flavivirdae is an important family of human and animal RNA viral pathogens (Rice, CM. 1996. Flavivirdae: The viruses and their replication. In: Fields BN, Knipe DM, Howley PM., eds. Fields virology. Philadelphia: Lippincott-Raven Publishers. pp. 931-960.) The three currently recognized genera of the Flavivirdae family exhibit distinct differences in 20 transmission, host range, and pathogenesis. For example, members of the classical flavivirus genus, such as yellow fever virus and dengue virus, are typically transmitted to vertebrate hosts via arthropod vectors and cause acute self-limiting disease (Monath TP, Heinz FX. 1996. Flaviviruses. In: Fields BN, Knipe DM, Howley PM., eds. Fields virology. New York: Raven Press. pp. 961-1034). The pestiviruses, such as bovine viral diarrhea virus (BVDV) 25 and classical swine fever virus (CSFV), cause economically important livestock disease and are spread by direct contact or the fecal-oral route (Thiel et al., 1996. Pestiviruses. In: Fields BN, Knipe DM, Howley PM., eds. Fields virology. New York: Raven Press. pp. 1059-1073). The most recently characterized Flavivirdae genus is the hepacivirus genus, the sole member of which is the common and exclusively human pathogen, hepatitis C virus (HCV). HCV is 30

transmitted by contaminated blood or blood products and is the most common agent of non-A, non-B hepatitis, affecting more that 1% of the population worldwide (Houghton, 1996. Hepatitis C viruses. In: Fields BN, Knipe DM, Howley PM., eds. *Fields virology*. Philadelphia: Lippincott-Raven Publishers. pp. 1035-1058.). Unlike flavivirus and pestivirus infections, which are usually eliminated by host immune response, chronic HCV infections are common and can cause mild to severe liver disease including cancer.

Despite these differences, members of the *Flavivirdae* family share common structural features and gene expression strategies. Virus particles consist of a lipid bilayer envelope with embedded transmembrane glycoproteins surrounding a protein-RNA nucleocapsid. Genome RNAs are single-stranded of positive polarity, and function as the sole mRNA species for translation of a single long open reading frame (ORF). This ORF is translated into a polyprotein which is processed by cellular and viral proteases into mature viral proteins. Structural proteins destined for incorporation into virus particles are encoded in the N-terminal portion of the polyprotein, while the nonstructural proteins which form components of the viral RNA replicase are encoded in the remainder.

Replication of the *Flavivirdae* RNA genome occurs via synthesis of a full-length negative-strand intermediate and is asymmetric, favoring synthesis of positive-strand RNAs. However, little is known about the details of this process. For all three genera of the *Flavivirdae* family, full-length functional cDNA clones have been constructed and RNAs transcribed from these cDNA templates are infectious. For flaviviruses and pestiviruses, mutagenesis of these clones and efficient RNA transfection of permissive cell cultures provides a means of probing the role of *cis* RNA elements and viral proteins in replicase assembly and function. Such analyses are not yet possible for HCV since this virus is unable to replicate efficiently in cell culture.

Like many other RNA viruses, it is believed the 5' and 3' terminal sequences of the Flavivirdae contain conserved cis-elements important for translation, RNA replication, and packaging (Bukh et al., Proc. Natl. Acad. Sci. USA 89:4942-4946, 1992; Deng et al., Nucleic Acids Res. 21:1949-1957, 1993; Cahour et al., Virol. 207:68-76, 1995; Kolykhalov et al., J. Virol. 70:3363-3371, 1996; Men et al., J. Virol. 70:3930-3937, 1996; Tanaka et al., J. Virol. 70:3307-3312, 1996; Huang HV. 1997. Evolution of the alphavirus promoter and the cisacting sequences of RNA viruses. In: Saluzzo J-F, Dodet B. eds. Factors in the emergence of arbovirus disesases. Paris: Elsevier Press, pp. 65-79; Mandl et al., J. Virol. 72:2132-2140, 1998). The 5' nontranslated region (NTR) functions initially at the level of translation. Similar to most cellular mRNAs, flavivirus genome RNAs are translated in a cap-dependent manner. These RNAs contain a 5' cap structure that is presumably added by virus-encoded

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RNA triphosphatases, guanylyl-, and methyl-transferases (Rice, 1996, *supra*). In contrast, the translational strategy employed by pestiviruses and HCV is more similar to that of the picornaviruses. These RNAs appear to be uncapped and contain long 5' NTRs with *cis* RNA elements that function as internal ribosome entry sites (IRES) for translation initiation at the polyprotein AUG (Lemon et al., *Semin. Virol.* 8:274-288, 1997).

The 5' NTRs of HCV and BVDV have a similar structural and functional organization despite containing only short stretches of high sequence identity (Wang et al., Curr. Top. Microbiol Immunol. 203:99-115, 1995; Lemon et al., 1997, supra). The IRES within each NTR is located at the 3' end of the NTR at a position proximal to the AUG initiation codon of the ORF. Although the 5' terminal sequence of each of these viruses is apparently not required for IRES function (Rijnbrand et al., FEBS Lett 365:115-119, 1995; Honda et al., Virology. 222:31-42, 1996; Rijnbrand et al., J. Virol. 71:451-457, 1997), these sequences are highly conserved among different strains of HCV (Bukh et al., Proc. Natl. Acad. Sci. USA:89:4942-4946, 1992) or BVDV (Deng et al., 1993, supra), suggesting they play other roles in viral replication. For example, sequences in the 5' NTR may be required for regulating translation versus initiation of negative-strand RNA synthesis. Such regulation could occur by direct interaction of 5' and 3' RNA elements or indirectly, via RNA-protein interactions. Sequences in the 5' NTR may also modulate packaging versus translation. Finally, sequences complementary to the 5' NTR, which are located at the 3' end of negative-strand RNA, are likely to function in the initiation of positive-strand RNA synthesis.

The HCV 3' NTR contains an internal polypyrimidine tract followed by a highly conserved sequence of 98 bases at the 3' terminus, which has been shown to be required for replication of HCV (U.S. Application Serial No. 08/811,566).

Further elucidation of the role of sequences in the HCV 5' and 3' NTRs has been hampered by the inefficient replication of HCV in cell culture. This aspect of HCV biology also makes it difficult to identify and test possible antiviral compounds for activity against HCV. Thus, a need exists for a system which facilitates investigation of HCV replication and therapeutic approaches to control HCV infections.

30 Summary of the Invention

Briefly, therefore, the present invention provides novel compositions and methods for studying HCV replication which are based on the discovery that chimeras of HCV and BVDV genomic sequences can be constructed that are able to replicate in cell culture. The BVDV-specific sequence provides the chimeric viral nucleic acid with the ability to replicate in cell culture, while the HCV-specific sequence allows the chimeric viral nucleic acid to be used to

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screen possible compounds for anti-viral activity against HCV. It is believed that similar replication-competent chimeras can be constructed from HCV and other pestiviruses.

Thus, in one embodiment, the present invention provides a novel, chimeric viral RNA in which at least one of the 5' NTR; ORF and 3' NTR regions is chimeric and comprises a nucleotide sequence from the corresponding region of a pestivirus in operable linkage with a nucleotide sequence from the corresponding region of an hepatitis C virus (HCV). The chimeric viral RNA is replication-competent. In preferred embodiments, the pestivirus is BVDV.

In other embodiments, the invention provides a polynucleotide comprising a DNAdependent promoter operably linked to a cDNA of a chimeric viral RNA as described above and cells transiently transfected or stably transformed with the polynucleotide. In some embodiments the cDNA may encode a dominant selectable marker or an assayable reporter.

In yet another embodiment, the invention provides a method for identifying compounds having anti-HCV activity. The method comprises providing a first cell containing a chimeric viral nucleic acid derived from HCV and a pestivirus as described above and a second cell containing the pestivirus, and then comparing the replication efficiency of the chimeric viral nucleic acid in the presence and absence of a test compound to the replication efficiency of the pestivirus in the presence and absence of the test compound, wherein a greater reduction in compound-induced replication efficiency of the chimeric viral nucleic acid than the pestivirus indicates the compound has anti-HCV activity.

The invention also provides a genetically-engineered virus which comprises a chimeric viral nucleic acid derived from HCV and a pestivirus as described above. In one embodiment the genetically-engineered virus comprises virus particles containing at least one HCV structural protein and is useful in a vaccine against HCV. In another embodiment, the genetically-engineered virus is attenuated as compared to the pestivirus and is useful as a vaccine against the pestivirus.

In a still further embodiment, the invention provides a replication-competent BVDV vector expressing a heterologous sequence. The BVDV vector comprises the BVDV sequences encoding the BVDV replication machinery. In some embodiments, the replication-competent BVDV vector expresses an antigen and is useful as a vaccine.

Brief Description of the Drawings

Figure 1 is a schematic representation of the 5' NTRs of BVDV, HCV, and EMCV showing the position of the start codons of the ORF, and the boxes indicating the canonical IRES elements.

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Figure 2 shows a schematic representation of BVDV and HCV chimeras, plaque phenotypes, reticulocyte translation efficiencies relative to parental BVDV, specific infectivities in MDBK cells, titers at 24 and 48 h post-transfection (or 72 h, as indicated), and an indication of whether pseudorevertants arose with results from BVDV, 5HCV, BVDV+HCV, and BVDV+HCVdelB3 chimeras shown in Fig. 2A and results from BVDV+HCVdelB2B3, BVDV+HCVdelB1B2B3, BVDV+HCVdelB2B3H1, and BVDV+HCVdelB2B3H1H2 shown in Fig. 2B, where N.D. means not determined.

Figure 3 illustrates the *in vitro* translation efficiency of BVDV RNA or chimeras showing bar graphs of the amount of N^{pro}, the N-terminal protein in the BVDV ORF, expressed by the various constructs.

Figure 4 illustrates a schematic representation of EMCV chimeras, plaque phenotypes, reticulocyte translation efficiencies relative to parental BVDV, specific infectivities in MDBK cells, titers at 24 and 48 h post-transfection (or 72 h, as indicated), and an indication of whether pseudorevertants arose.

Figure 5 illustrates a pseudorevertant analyses showing in (Fig. 5A) the relative positions of mutations detected within the plaque-purified variants of passaged BVDV+HCVdelB1B2B3, 5'EMCV, and 5'HCV, and in (Fig. 5B) the 5' terminal sequences of pseudorevertants of BVDV+HCVdelB1B2B3, 5'EMCV, and 5'HCV. Novel nucleotides or sequences are shown in bold upper case type. Pseudorevertants are numbered and designated by the suffix ".R". The upper case sequence in BVDV+HCVdelB1B2B3 and BVDV+HCVdelB1B2B3.R1 is a remnant of downstream BVDV 5' NTR sequences and was created during the cloning procedures.

Figure 6 illustrates the construction of derivatives of 5'HCV designed to contain 5' termini corresponding to the sequence detected within the three analyzed pseudorevertants. Fig. 6A shows the 5' terminal sequence of the 5'HCV derivatives with the suffix (orig) designating a derivative containing the original 5' terminal sequence of the pseudorevertant; the suffix (cons) designating a derivative containing the consensus tetranucleotide sequence 5'-GUAU at the same position; and novel sequences shown in bold upper case type. Fig. 6B shows plaque phenotypes, reticulocyte translation efficiencies relative to parental BVDV, specific infectivities in MDBK cells, and titers at 24 and 48 h post-transfection are indicated.

Figure 7 illustrates a single step growth curve for various chimeric constructs showing released virus titers measured by performing plaque assays on MDBK cells transfected with various constructs.

Figure 8 illustrates replication of BVDV RNA or chimeric derivatives in transfected

35 MDBK cells. Equal numbers of MDBK cells (~ 8 x 10⁶) were electroporated with 5 \(\precedge \) g of

each in vitro synthesized RNA. MDBK cells were also transfected with infectious yellow fever 17D and Sindbis RNAs to provide molecular mass markers. One fifth of the transfected cells were seeded on 35-mm dishes and incubated in D-MEM supplemented with 10% horse serum for 6 h at 37°C. The media were then replaced with 1 ml of fresh media containing 2 g/ml of actinomycin D and 40 Ci/ml of ³H-uridine. Incubations were continued for 10 h at 37°C. RNAs were isolated as described in Materials and Methods, and 1/4 of the samples was denatured in glyoxal and loaded on an agarose gel. (A) Autoradiograph of the dried gel. Only the portion of the gel containing the genomic RNAs is shown. (B) Amount of radioactivity contained within the displayed fragments as determined by scintillation counting. BVDV, lane 1; 5'HCV, lane 2; BVDV+HCVdelB2B3, lane 3; BVDV+HCVdelB2B3H1, lane 4; 5'HCV.R1orig, lane 5; 5'HCV.R1cons, lane 6; 5'HCV.R3orig, lane 7; 5'HCV.R3cons, lane 8; 5'HCV.R2corig, lane 9; 5'HCV.R2cons, lane 10; yellow fever 17D, lane 11; Sindbis, lane 12; non-transfected MDBK cells, lane 13. The

experiments shown is one of two repetitions which yielded similar results.

Figure 9 illustrates the genetic map of plasmid pACNR/BUD.

Figure 10 illustrates the sequence of low copy number plasmid pACNR/BVDV NADL (circular) harboring the functional cDNA of cytopathic BVDV NADL (positive sense cDNA 5' to 3'; nt 1-12578.

Figure 11 illustrates the sequence of infectious BVDV NADL (positive sense cDNA 20 5' to 3').

Figure 12 illustrates the sequence of infectious non-cytopathic BVDV NADL lacking cIns (positive sense cDNA 5' to 3').

Figure 13 illustrates the sequence adapted HCV 5' NTR from 5'HCV/R1.cons (positive sense cDNA 5' to 3'; only the sequence from the 5' base to the ATG initiating the polyprotein is shown).

Figure 14 illustrates the sequence of adapted HCV 5' NTR from 5'HCV/R1.orig (positive sense cDNA 5' to 3'; only the sequence from the 5' base to the ATG initiating the polyprotein is shown).

Figure 15 illustrates the sequence of adapted HCV 5'NTR from 5'HCV/R2.cons (positive sense cDNA 5' to 3'; only the sequence from the 5' base to the ATG initiating the polyprotein is shown).

Figure 16 illustrates the sequence of adapted HCV 5' NTR from 5'HCV/R2.orig (positive sense cNDA 5' to 3'; only the sequence from the 5' base to the ATG initiating the polyprotein is shown).

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Figure 17 illustrates the sequence of adapted HCV 5' NTR from 5'HCV/R3.cons (positive sense cDNA 5' to 3'; only the sequence from the 5'base to the ATG initiating the polyprotein is shown).

Figure 18 illustrates the sequence of adapted HCV 5'NTR from 5'HCV/R3.orig (positive sense cDNA 5' to 3'; only the sequence from the 5' base to the ATG initiating the polyprotein is shown).

Figure 19 illustrates the sequence of prototype HCV-BVDV chimera from pNADL/5'HR3.orig/3'H3'B with the adapted HCV 5'NTR from 5'HCV/R3.orig and tandem 3' NTR elements from HCV followed by BVDV (positive sense cDNA 5' to 3') as discussed in Example 5.

Figure 20 illustrates various deletions of the poly U track in the 3'NTR HCV sequence of BVDV/HCV chimera p5H-3H33.

Figure 21 illustrates the schematic representation of functional HCV/-BVDV chimera from pCBV/p7.

Figure 22 illustrates the sequence of functional HCV-BVDV chimera from pCBV/p7 (positive sense cDNA 5' to 3').

Figure 23 illustrates the schematic representation of a HCV/BVDV chimera with selectable marker.

Figure 24 illustrates the sequence of functional HCV-BVDV chimera from pCBV/p7/IRES-pac expressing a dominant selectable marker conferring resistance to puromycin (positive sense cDNA 5' to 3').

Figure 25 illustrates the schematic representation of a bicistronic HCV/BVDV chimera.

Figure 26 illustrates the sequence of functional bicistronic chimera expressing the
entire HCV structural region derived from plasmid pNADL/BI#41/HCV str (positive sense cDNA 5' to 3')

Description of the Preferred Embodiments

In accordance with the present invention, the inventors herein have succeeded in generating HCV-BVDV chimeric RNAs which are replication competent. Such chimeras are useful in screening compounds *in vitro* for antiviral activity against HCV. In addition, it is believed that *in vivo* replication of HCV-BVDV chimeras according to the invention may be attenuated as compared to wild-type BVDV and thus may be useful in vaccinating animals against BVDV. It is also believed that the HCV chimeric structures described herein for BVDV are applicable to other pestiviruses.

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In the context of this disclosure, the following terms will be defined as follows unless otherwise indicated:

"Cis-acting sequences" means the nucleotide sequences from an RNA virus genome that are necessary for recognition of the genomic RNA by specific protein(s) of the RNA virus or host cell that carry out replication, transcription, translation or packaging of the genome.

"Genetically-engineered virus" means any virus whose genome is different than that of a wild-type virus due to a human-made deletion, insertion, or substitution of one or more nucleotides to the wild-type viral genome.

"Infectious" when used to describe a virus means the virus is capable of entering cells and initiating a virus replication cycle, whether or not this leads to the production of new RNA virus particles.

"Nucleotide sequence" as used herein refers to DNA and the corresponding RNA sequence where relevant. It will be understood that sequences shown in the Figures are DNA versions of the RNA sequence and that chimeric molecules of the invention may comprises RNA molecules or cDNA copies of such RNA molecules.

"Replication-competent" as applied to a chimeric HCV-pestivirus RNA means the RNA is capable of RNA-dependent replication in at least one cell type that supports replication of the wild-type parental pestivirus. The number of replicated RNA molecules produced by an HCV-pestivirus chimeric RNA of the invention is at least 10-fold higher than the limit of detection, which is typically 10 to 100 molecules. More preferably, chimeric RNA production by the HCV-pestivirus chimeric RNA is at least 10² to 10³-fold higher than the detection limit. The replication-competent chimeric RNA replicates at an efficiency that is preferably, at least 0.001%, more preferably, at least 0.11%, more preferably at least 10% and most preferably at least 50% up to 90% that of the parental pestivirus in the same cell type.

"Transfected cell" means a cell containing an exogenously introduced nucleic acid molecule, and includes cells that are transiently transfected with the exogenous nucleic acid.

"Transformed cell" or "stably transformed cell" means a cell containing an exogenously introduced nucleic acid molecule which is present in the cytoplasm or nucleus of the cell and may be stably integrated into the chromosomal DNA of the cell.

"Virus" means a virion, virus particle or a viral genome.

A chimeric viral RNA according to the invention is designed to comprise a 5' NTR, an ORF, and a 3' NTR, at least one of which is a chimeric region containing two operably linked nucleotide sequences that are from the same region of a pestivirus and an HCV.

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Pestivirus-specific sequences useful in the invention can be taken from the appropriate genomic region of any cytopathic or noncytopathic type I or type II BVDV isolate, classical swine fever virus (CSFV) isolate, or border disease viral isolate. For a list of pestiviruses, see Thiel, H.-J., P. G. W. Plagemann, and V. Moennig. 1996. Pestiviruses, p. 1059-1073. In B. N. Fields, D. M. Knipe and P. M. Howley (ed.), Fields Virology. Raven Press, New York. HCV-specific sequences can be taken from any strain or isolate of HCV, including but not limited to HCV-1, HCV-1a, HCV-1b, HCV-1c, HCV-2a, HCV-2b, HCV-2c, HCV-3a. Preferably, the parental pestivirus is a cytopathic strain of BVDV and the parental HCV strain is HCV-1.

The pestivirus- and HCV-specific sequences are operably linked in the chimeric region, meaning the sequences are arranged such that the resulting chimeric structure is functional in the context of replication of the pestivirus. For example, in one preferred embodiment the chimeric viral RNA comprises a chimeric 5' NTR which comprises a BVDV-specific 5' terminal sequence of 5'-(G/A)UAU and an IRES derived from HCV, with the ORF and the 3' NTR consisting of a sequence from the same regions of BVDV. The BVDV-specific sequences at the 5' terminus and in the ORF and 3' NTR are chosen such that they are functional in the context of BVDV, meaning the chimeric viral RNA expresses the replication machinery of BVDV and this replication machinery is capable of replicating the chimeric RNA. In addition, translation of the BVDV ORF in the chimeric viral RNA is dependent upon a functional HCV IRES. The presence of a functional HCV IRES in this chimera allows the chimera to be used to screen for compounds that target the HCV IRES and thereby inhibit translation of the BVDV ORF as well as replication of the chimeric virus. Such compounds would be expected to also inhibit translation of the ORF in a wild-type HCV and consequently inhibit HCV replication.

Compounds that could be screened for anti-HCV activity using this and other HCV-BVDV 5' NTR chimeras include but are not limited to antisense RNAs, RNA decoys that bind proteins involved in recognition of the HCV-specific sequences, ribozymes, and small molecule inhibitors of critical RNA-protein interactions. The use of such substances for therapeutic applications are known in the art. See, e.g., Amarzguioui M, et al., "Hammerhead ribozyme design and application." *Cell Mol Life Sci.* 1998 Nov;54(11):1175-202; Welch PJ, et al., "Expression of ribozymes in gene transfer systems to modulate target RNA levels.", *Curr Opin Biotechnol.* 1998 Oct;9(5):486-96; Bramlage B, et al. "Designing ribozymes for the inhibition of gene expression."; *Trends Biotechnol.* 1998 Oct;16(10):434-8; Gewirtz AM, et al. "Nucleic acid therapeutics: state of the art and future prospects."; *Blood.* 1998 Aug 1:92(3):712-36; Altman S., "RNase P in research and therapy." *Biotechnology* (N Y). 1995

Apr; 13(4):327-9; Flanagan WM., "Antisense comes of age."; Cancer Metastasis Rev. 1998 Jun; 17(2):169-76; Agrawal S, et al., "Antisense therapeutics." Curr Opin Chem Biol. 1998 Aug;2(4):519-28; Caselmann WH, et al., "Synthetic antisense oligodeoxynucleotides as potential drugs against hepatitis C." Intervirology 1997;40(5-6):394-9; Neckers LM., 5 "Oligodeoxynucleotide inhibitors of function: mRNA and protein interactions." Cancer J Sci Am. 1998 May; 4 Suppl 1:S35-42; Agrawal S, et al. "Mixed backbone oligonucleotides: improvement in oligonucleotide-induced toxicity in vivo." Antisense Nucleic Acid Drug Dev. 1998 Apr;8(2):135-9; Crooke ST., "An overview of progress in antisense therapeutics." Antisense Nucleic Acid Drug Dev. 1998 Apr;8(2):115-22; Fraisier C, et al., "High level. 10 inhibition of HIV replication with combination RNA decoys expressed from an HIV-Tat inducible vector."; Gene Ther. 1998 Dec;5(12):1665-76; Gervaix A, et al. "Gene therapy targeting peripheral blood CD34+ hematopoietic stem cells of HIV-infected individuals." Hum Gene Ther. 1997 Dec 10;8(18):2229-38; Nakaya T, et al. "Inhibition of HIV-1 replication by targeting the Rev protein." Leukemia 1997 Apr;11 Suppl 3:134-7; Nakaya T, et 15 al. "Decoy approach using RNA-DNA chimera oligonucleotides to inhibit the regulatory function of human immunodeficiency virus type 1 Rev protein." Antimicrob Agents Chemother. 1997 Feb;41(2):319-25; Smith C, et al. "Transient protection of human T-cells from human immunodeficiency virus type 1 infection by transduction with adeno-associated viral vectors which express RNA decoys." Antiviral Res. 1996 Oct;32(2):99-115; Bahner I, et 20 al. "Transduction of human CD34+ hematopoietic progenitor cells by a retroviral vector expressing an RRE decoy inhibits human immunodeficiency virus type 1 replication in myelomonocytic cells produced in long-term culture." J Virol. 1996 Jul;70(7):4352-60; Lee SW, et al. "Inhibition of human immunodeficiency virus type 1 in human T cells by a potent

Rev response element decoy consisting of the 13-nucleotide minimal Rev-binding domain." J Virol. 1994 Dec;68(12):8254-64; Lisziewicz J, et al. "Inhibition of human immunodeficiency virus type 1 replication by regulated expression of a polymeric Tat activation response RNA decoy as a strategy for gene therapy in AIDS." Proc Natl Acad Sci U S A. 1993 Sep 1;90(17):8000-4; Bevec D, et al. "Inhibition of human immunodeficiency virus type 1 replication in human T cells by retroviral-mediated gene transfer of a dominant-negative Rev trans-activator." Proc Natl Acad Sci U S A. 1992 Oct 15;89(20):9870-4.

It is contemplated that a number of replication-competent chimeric structures can be made that allow the function of various HCV sequence elements and proteins to be studied and targeted in drug screening assays. For example, the invention includes replication-competent HCV-pestivirus chimeras having a chimeric ORF. One such chimeric ORF is one comprising an HCV sequence encoding the structural proteins and a pestivirus sequence

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encoding the nonstructural proteins. It is believed that upon introduction into a cell, such a HCV-BVDV ORF chimera will produce HCV-like virus particles that will be released from the cell and capable of infecting cells normally infected by wild-type HCV, i.e., cells expressing an HCV receptor such as human CD81. Such ORF chimeras would be useful to screen compounds for drugs that inhibit formation, release or entry of HCV particles. In addition, ORF chimeras that produce virus particles containing at least one HCV structural protein would be useful as vaccines against HCV. Other ORF chimeras contemplated by the invention include, for example, chimeras comprising a pestivirus sequence encoding structural proteins and an HCV sequence encoding one or more nonstructural proteins such as the NS3 protease, NS4A cofactor, NS5A phosphoprotein/interferon resistance determinant and/or the NS5B polymerase. Replication of such ORF chimeras would be dependent upon the function of the HCV nonstructural protein(s) and these ORF chimeras could be used to screen for drugs that target the HCV nonstructural protein(s) as well as to screen for and map potential drug resistance mutations in HCV nonstructural proteins. In addition, HCVpestivirus ORF chimeras could be useful for developing alternative in vivo animal models for HCV replication and HCV-associated hepatocellular carcinoma to evaluate antivirals and anti-tumor agents.

The invention also provides replication-competent HCV-pestivirus chimeras having a chimeric 3' NTR which contains one or more conserved elements of the HCV 3' NTR. Such 3' NTR chimeras would be useful for screening or evaluating compounds targeted against the HCV 3' NTR. Compounds that could be screened include antisense RNA molecules, ribozymes and small molecule inhibitors of critical RNA-protein interactions. One 3' NTR chimera according to the invention comprises a BVDV 5' NTR, BVDV ORF and a chimeric 3' NTR which consists of an HCV-specific sequence derived from the HCV 3' NTR immediately followed by a BVDV 3' NTR. The HCV-specific 3' NTR that allows for replication in the context of BVDV has a deletion in the 3' NTR poly (U) tract but has all the other HCV 3' NTR elements, including the 98 bp 3' terminal conserved element.

HCV-pestivirus chimeras included within the scope of the invention include those comprising combinations of chimeric regions, i.e., 5' NTR and ORF chimeras; 5' NTR and 3' NTR chimeras; ORF and 3' NTR chimeras; and chimeric RNAs in which each of the 5' NTR, ORF and 3' NTR regions comprise an HCV sequence operably linked to a pestivirus sequence.

The invention also provides chimeric RNAs having two ORFs, or bicistronic HCV-pestivirus chimeras. Bicistronic chimeras contemplated by the invention include structures in which the first ORF contains one or more HCV genes and is followed by a second IRES

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operably linked to a second ORF encoding the pestivirus replicase machinery. It is also contemplated the first ORF may encode a heterologous sequence such as an antigen.

It is believed that many HCV-pestivirus chimeras of the invention will be attenuated as compared to the parental wild-type pestivirus. Such attenuated chimeric RNA genomes would be candidate vaccines in the form of live-attenuated virus particles or as RNA or cDNA "genetic" vaccines.

The invention also includes vaccines against HCV which comprise an immunogenically-effective amount of HCV-pestivirus particles or nucleic acid. Anti-HCV vaccines comprising virus particles should preferably contain one or more HCV structural proteins.

The therapeutic or pharmaceutical compositions of the present invention can be administered by any suitable route known in the art including for example by injection such as intraperitoneal, intravenous, subcutaneous, intramuscular, transdermal, intrathecal or intracerebral injection. Administration can be either rapid as by injection or over a period of time as by slow infusion or administration of slow release formulation.

Compositions according to the invention can be employed in the form of pharmaceutical or veterinary preparations. Such preparations are made in a manner well known in the pharmaceutical and veterinary arts. One preferred preparation utilizes a vehicle of physiological saline solution, but it is contemplated that other pharmaceutically acceptable carriers such as physiological concentrations of other non-toxic salts, five percent aqueous glucose solution, sterile water or the like may also be used. It may also be desirable that a suitable buffer be present in the composition. Such solutions can, if desired, be lyophilized and stored in a sterile ampoule ready for reconstitution by the addition of sterile water for ready injection. The primary solvent can be aqueous or alternatively non-aqueous.

The carrier can also contain other pharmaceutically-acceptable excipients for modifying or maintaining the pH, osmolarity, viscosity, clarity, color, sterility, stability, rate of dissolution, or odor of the formulation. Similarly, the carrier may contain still other pharmaceutically-acceptable excipients for modifying or maintaining release or absorption or penetration across the blood-brain barrier. Such excipients are those substances usually and customarily employed to formulate dosages for parenteral administration in either unit dosage or multi-dose form or for direct infusion into the cerebrospinal fluid by continuous or periodic infusion.

It is also contemplated that certain formulations containing a chimeric virus according to the invention are to be administered orally. Such formulations are preferably encapsulated and formulated with suitable carriers in solid dosage forms. Some examples of suitable

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carriers, excipients, and diluents include lactose, dextrose, sucrose, sorbitol, mannitol, starches, gum acacia, calcium phosphate, alginates, calcium silicate, microcrystalline cellulose, polyvinylpyrrolidone, cellulose, gelatin, syrup, methyl cellulose, methyl- and propylhydroxybenzoates, talc, magnesium, stearate, water, mineral oil, and the like. The formulations can additionally include lubricating agents, wetting agents, emulsifying and suspending agents, preserving agents, sweetening agents or flavoring agents. The compositions may be formulated so as to provide rapid, sustained, or delayed release of the active ingredients after administration to the patient by employing procedures well known in the art. The formulations can also contain substances that diminish proteolytic degradation and promote absorption such as, for example, surface active agents.

The specific dose is calculated according to the approximate body weight or body surface area of the patient or the volume of body space to be occupied. The dose will also be calculated dependent upon the particular route of administration selected. Such calculations can be made without undue experimentation by one skilled in the art. Exact dosages are determined in conjunction with standard dose-response studies. It will be understood that the amount of the composition actually administered will be determined by a practitioner, in the light of the relevant circumstances including the condition or conditions to be treated, the choice of composition to be administered, the age, weight, and response of the individual patient, the severity of the patient's symptoms, and the chosen route of administration. Dose administration can be repeated depending upon the pharmacokinetic parameters of the dosage formulation and the route of administration used.

Replication-competent HCV-pestiviruses are generated by choosing the HCV function or sequence element desired to be studied. The HCV sequence can be obtained from a plasmid clone of a partial or full HCV genome using PCR to amplify a target region containing the desired sequence or by restriction enzyme digestion. The HCV fragment is then inserted into the desired location of a clone of the pestivirus genome using standard techniques. Desired portions of the pestivirus genome may be deleted before or after addition of the HCV fragment. The recombinant genome is then transfected into a cell that supports replication of the parental pestivirus genome and their ability to replicate using standard assays. For example, replication can be assessed by virus-induced cytopathic effect; plaque formation; detection of viral antigens and/or viral RNA accumulation; and by plaque assay measuring released infectious virus. The inventors herein have found that the BVDV RNA replication machinery works in many cell types, including bovine, hamster, mouse and human cells. It has also been reported that BVDV RNAs can amplify in other cell types including human hepatoma lines and hepatocytes (Behrens SE, et al., J Virol. 1998 Mar;72(3):2364-72).

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The host cell range for a particular chimera will be dependent upon the properties of that chimera as empirically determined.

As described below, some chimeras do not replicate stably as indicated by heterogeneity in the size of plaques produced by the chimeric virus. Upon passage, pseudorevertants can frequently be isolated that are capable of stable replication. Such pseudorevertants will have one or more deletions or base substitutions in the HCV and/or pestivirus sequences. Information derived from these gain-of-function mutations can be used to define the elements necessary for generating stable, replication-competent chimeras of HCV and a pestivirus.

The invention provides a method for screening compounds for antiviral activity against HCV. The method involves comparing a test compound's effect on replication of a chimeric HCV-pestivirus RNA molecule as described above with the compound's effect on replication of the parental pestivirus. Compounds which have a greater effect on replication of the chimeric virus than the pestivirus are likely directed against the HCV portion of the chimera. Typically, the method is performed by providing duplicate cell cultures containing a chimeric viral RNA which is replication-competent in that cell, treating one of the culture with the test compound, and then measuring the replication efficiency of the chimeric RNA in both cultures. Any effect induced by the compound is compared against the compound's effect on replication of the parental pestivirus in cells of the same type. This control assay is preferably performed at the same time using the same culture conditions.

The cells used in the screening assay can be prepared by transfertly transfecting the cells with the desired chimeric RNA molecule as described below. Alternatively, it is contemplated that the chimeric RNA molecule can be constitutively expressed in the cell by transfecting the cell with a polynucleotide comprising a cDNA of the chimeric RNA operably linked to a DNA-dependent promoter. The chimeric cDNA may include a selectable marker. which would allow for selection of cells expressing the chimeric RNA. It is also envisioned the selectable marker could be a dominant marker that allows selection of cells expressing chimeras having adaptive mutations or selection of cells permissive for virus replication (Frolov et al., *J. Virol.* 73:3854-3865, 1999). It is also contemplated the cDNA could express a reporter gene that could be assayed to measure RNA replication.

Alternatively, chimeric virus particles are incubated with a cell permissive for infection by the pestivirus in the presence or absence of the test compound and then replication of the chimeric virus is measured and compared to the replication of the parental pestivirus incubated with the same cell type in the presence or absence of the test compound.

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Inhibition of replication can be measured in many ways, including assaying for the reduction of virus-induced cytopathic effect; inhibition of plaque formation, reduced production of viral antigens as detected by immunofluoresence assay; reduced viral RNA accumulation; reduction in released infectious virus from treated and untreated control and chimera samples using a plaque assay. In addition, it is contemplated that a cell line that is designed for pestivirus-specific transactivation of a reporter gene could be used directly or in lieu of a plaque assay. The reporter gene is operably linked to a promoter that is activated upon infection by the chimeric virus and production of the viral transactivator protein.

Preferred embodiments of the invention are described in the following examples.

Other embodiments within the scope of the claims herein will be apparent to one skilled in the art from consideration of the specification or practice of the invention as disclosed herein. It is intended that the specification, together with the examples, be considered exemplary only, with the scope and spirit of the invention being indicated by the claims which follow the examples.

15 Example 1

This example illustrates the construction and analysis of 5' HCV-BVDV chimeras as reported in detail in Frolov et al. (RNA 4:1418-1435, 1998) which is incorporated in its entirety by reference. A functional clone of BVDV (Mendez et al., J. Virol. 72:4737-4745, 1998) was used to construct and characterize a series of 5' NTR chimeras with sequences derived from HCV and the picornavirus, encephalomyocarditis virus (EMCV). The results help to define the requirements of a functional BVDV 5' NTR and provide replication-competent BVDV-HCV chimeras dependent on a functional HCV IRES.

Example 2

This example illustrates the construction of chimeras for expressing additional functional portions of the HCV genome by addition of further HCV sequence downstream from the functional or adapted HCV 5'NTR chimeras fused in-frame to the BVDV ORF.

One such construct (Figure 21) involves fusion of HCV sequences to BVDV sequences in the p7 protein coding region (at a convenient BseRI restriction site). Both HCV and BVDV encode a p7 protein that is located immediately downstream of the E2 protein. The p7 protein is a small hydrophobic protein of unknown function. pCBV/p7 consists of the first 79 bases of the BVDV 5'NTR encoding stem loop structure B1' and B1, followed by the entire HCV 5'NTR, the entire HCV structural protein coding region and the first 36 amino acids of HCV p7 fused to the C-terminal 31 amino acids of BVDV p7. The fused p7 gene is followed by the remainder of the BVDV ORF including the entire nonstructural region and the BVDV 3' NTR. Transfection of MDBK cells with the RNA corresponding to this

sequence (Figure 22) leads to replication of the chimeric RNA and production of the expected HCV and BVDV polyprotein cleavage products. Variations on this strategy are envisioned in which all or part of the HCV polyprotein and cis elements important for RNA packaging can be expressed in viable chimeras. In addition the BVDV replicase regions for either cytopathic or non-cytopathic pestiviruses (like NADL clns-) can be used. Transfection of cells permissive for HCV particle, assembly, release and reinfection with this chimeric RNA can be used to make HCV-like particles. These particles and this infection system can be used (i) to screen for specific inhibitors of HCV particle, assembly, release and reinfection, (ii) for identifying antibodies capable of neutralizing HCV infectivity and (iii) as live or inactivated vaccines. Furthermore, this embodiment of the invention demonstrates that the BVDV RNA replication machinery can be used for expression of heterologous RNA and polypeptide sequences and can be used as a vehicle for RNA or DNA "genetic" vaccination in which the BVDV replicase amplifies the level of antigen expression by cytoplasmic RNA-dependent replication.

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Example 3

This example illustrates chimeric RNA's that are modified to express dominant selectable markers, assayable markers or FACS sortable markers.

Such variants can be used to select for chimeras capable of replication in particular cell types, or to screen for cell types that are permissive for replication of the chimeric RNA. Selectable markers include, but are not limited to, the genes encoding puromycin resistance (puromycin N-acetyl transferase; PAC), neomycin resistance, blasticidin resistance, hygromycin resistance, etc. Assayable markers include, but are not limited to, the genes encoding B-galactosidase, luciferase, B-glucuronidase, etc. Easily sortable molecules include single chain antibodies, cell surface markers, and non-toxic protein markers like green fluorescent protein. In a specific example (Figures 23 and 24), the RNA encoded by pCBV/p7 was modified to include a cassette at the beginning of the BVDV 3'NTR that is comprised of the EMCV IRES driving the gene encoding PAC. This chimeric RNA can replicate, expresses PAC and confers resistance to puromycin resistance. This property can be used to select for variants of the chimera that are capable of noncytopathic replication in desired cells type and also provides a means of showing that cells harbor a functional chimeric RNA. Desired variants can be identified, cloned and further characterized as described in Example 1. Of note, is that this location in the BVDV genome and this strategy for expressing heterologous genes may also be applied to using infectious attenuated

pestiviruses as gene expression vectors and as chimeric live vaccines against other animal pathogens.

Example 4

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This example illustrates the use of the bicistronic strategy as an alternative to the inframe fusions described in Example 2.

A specific example is shown in Figure 25 and its sequence as Figure 26. In this bicistronic chimera, the 5' sequences are identical to that of pCBV/p7 except that the HCV ORF continues to include the first 246 amino acids of NS4B. The HCV sequence is followed by the EMCV IRES fused to BVDV Npro, the N-terminal 10 aa of BVDV C, the C-terminal 19 aa of C, 9 N-terminal amino acids of Erns, 48 C-terminal amino acids of E2 and the remainder of the BVDV NADL ORF and 3' NTR. The constructed BVDV ORF encodes a functional BVDV RNA replicase. The deletions in the N-terminal portion of this ORF were designed to preserve proper membrane topology and processing of the replicase. The bicistronic chimeric RNA can replicate upon transfection of permissive BVDV host cells.

Example 5

This example illustrates 3'NTR chimeras. Although initial attempts to recover viable chimeric viruses in which the BVDV 3'NTR was completely replaced by that of HCV were unsuccessful, a strategy similar to that detailed in Example 1 has produced chimeras that harbor the conserved elements of the HCV 3'NTR. An initial tandem 3'NTR construct was made in which the HCV 3'NTR was engineered to follow the BVDV ORF. The complete BVDV 3'NTR was position 3' to the HCV 3' NTR after a short heterologous sequence. This sequence of this parental construct, which replicated poorly, is shown in Figure 19 RNAs transcribed from this plasmid were of low specific infectivity suggesting that revertants or pseudorevertants might have arisen. Indeed isolation and sequence analysis of several independent plaque-forming variants revealed that deletions in the HCV poly U tract of various lengths had occurred. These revertant sequences are shown in Figure 20. When these altered HCV 3'NTRs were reconstituted into the original tandem 3' NTR parent, they gave rise to plaque forming RNA transcripts of high specific infectivity, demonstrating that these alterations restored the ability of the chimeric RNA to replicate. Large deletions in the U tract gave rise to virus with more robust replication and larger plaques while stably maintaining the conserved HCV 3'NTR 98-base element and the polypyrimidine "transition" region. Such

chimeric viruses can now be used to screen and evaluate antisense, ribozyme, and other therapeutics targeted against this conserved HCV RNA element that is essential for replication.

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Materials and Methods

Plasmid Constructs

pACNR/BVDV NADL was previously described (Mendez et al., 1998, supra).

pBVDV is a derivative of pACNR/BVDV NADL which contains a G→T transversion at nt 14994 that creates an Xba I site upstream of the T7 promoter (T. Myers & C.M. Rice, unpubl.). To facilitate construction of the chimeras, subclones were created. First, two fragments were isolated by PCR amplification of p90/HCVFLIongpU (Kolykhalov et al., Science 277:570-574, 1997) with primers #498 (5'-TGTACATGGCACGTGCCAGCCCC) and #498 (5'-GATCAACTCCATGGTGCACGGTCT) and pBVDV with primers #481 (5'-AGACCGTGCACCATGGAGTTGATC) and #482 (5'-

- 15 CGTTTCACACATGGATCCCTCCTC). These two fragments were digested with ApaL I and ligated to produce a fragment containing a fusion of the HCV 5' NTR to the BVDV ORF. This fragment was digested with SacI and ligated into pGEM3Zf(-) which had been digested with Sma I and Sac I to produce the subclone pGEM498-Sacl. Next, a fragment containing the BVDV 5' NTR was synthesized by PCR amplification of pBVDV with primers #183 (5'-
- 20 TTTTCTAGATAATACGACTCACTATAGTATACGAGAATTAGAAAAGGCACTCG)
 and #480 (5'-GGGGGCTGGCACGTGCCATGTACA). This fragment was digested with

 Xba I and BsrG I and ligated into pGEM498-SacI digested with the same two enzymes, to
 create the plasmid pGEMXbal-SacI. pGemXbal-SacI contains a tandem fusion of the BVDV
 5' NTR, the HCV 5' NTR, and the 5' portion of the BVDV Npro gene. pBVDV + HCV was
 created by digesting pGEMXbal-SacI with Yba I and SacI and lighting the fragment into
- created by digesting pGEMXbal-SacI with Xba I and Sac I and ligating the fragment into pBVDV digested with the same two enzymes, and as such pBVDV + HCV contains the T7 promoter, followed by the entire 385-nt 5' NTR of BVDV, a GT dinucleotide (nt 386-387), the entire 341-nt 5' NTR of HCV (nt 388-728), and the sequence of the BVDV NADL strain including the ORF and 3' NTR. Derivatives of pBVDV + HCV containing deletions within the BVDV 5' NTR and/or the HCV 5' NTR were created in the subclone pGEMXbal-Sacl, as described below, prior to ligation into Sba I- and Sac I-digested pBVDV. For making deletions, restrictions sites with non-compatible protruding ends were treated with the Klenow fragment of DNA polymerase I prior to ligation. For creation of pBVDV +
- 35 BsrG I. For pBVDV + HCVdelB2B3 (deletion of nt 67-374), pGEMXbal-Sacl was digested

HCVdelB3 (deletion of nt 174-374, inclusive), pGEMXbal-Sacl was digested with Afl II and

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with Avr II and BsrG I. For pBVDV + HCVdelB1B2B3 (deletion of nt 33-374), pGEMXbal-Sacl was digested with SnaB I and BsrG I. For pBVDV + HCVdelB2B3H1 (deletion of nt 67-3396), pGEMXbal-Sacl was digested with Avr II and Xcm I. For pBVDV + HCVdelB2B3H1H2 (deletion of nt 67-513), pGEMXbal-Sacl was digested with AVR II and Bsg I. For pBVDV + HCVdelB2B3H3 (deletion of nt 67-374, 518-704), subclone pGEMXbal-SacidelB2B3 was digested with Sma I. p5'HCV was created by digesting p90/HCVliongpU with Xba I and Nru I and ligating the fragment into pBVDV + HCV digested with the same two enzymes.

The EMCV plasmid, pECg, was provided by Ann Palmenberg and is described elsewhere (Hahn et al., J. Virol 69:2697-2699, 1995). p5'EMCV contains the entire 710 nt of the 5' NTR of EMCV, followed by the open reading frame of BVDV and the 3' NTR. One extra G residue was added between the T7 promoter and the first nucleotide of the EMCV 5' NTR to facilitate efficient in vitro transcription. Convenient restriction sites within the BVDV 5' NTR or the EMCV 5' NTR were used to create additional chimeras. Sites with noncompatible protruding ends were treated with the Klenow fragment of DNA polymerase I prior to ligation. For example, the plasmid pBVDV + EMCVdelA contains nt 1-378 of BVDV 5' NTR fused with nt 45-710 of EMCV (the BsrG I site of BVDV ligated to the EcoR V site of EMCV), pBVDV + EMCVdelB3A contains nt 1-173 of BVDV fused with nt 45-710 of EMCV (the Afl II site of BVDV ligated to the EcoR V site of EMCV). pBVDV + EMCVdelB2B3A contains nt 1-66 of BVDV fused with nt 45-710 of EMCV (the Avr II site of BVDV ligated to the EcoR V site of EMCV). pBVDV + EMCVdelB3ABC contains nt 1-173 of BVDV fused with nt 161-710 of EMCV (the Afl II site of BVDV ligated to the Psp1405 site of EMCV). pBVDV + EMCVdelB2B3ABC nt 1-66 of BVDV fused with nt 161-710 of EMCV (the Avr II site of BVDV ligated to the Psp1406 site of EMCV). pBVDV + EMCVdelB3A-H contains nt 1-101 of BVDV fused with nt 289-710 of EMCV (the Nhe I 25 site of BVDV ligated to the Avr II site of EMCV). pBVDV + EMCVdelB2B3A-H contains nt 1-62 of BVDV fused with nt 289-710 of EMCV (the Avr II site of BVDV ligated to the Avr II site of EMCV). The schematics of the chimeric 5' NTRs are presented in Figures 2 and 4.

All other heterologous 5' NTRs used in the study were generated by PCR using an oligonucleotide complementary to nt256-272 of the HCV 5' NTR and primers containing the sequence of the Xba I restriction site followed by the T7 promoter, the heterologous sequences found in sequenced pseudorevertants, or sequences corresponding to different regions of the HCV 5' NTR. All the fragments were subcloned into the plasmid, pRS2 (a derivative of pUC19), sequenced, and recloned into the p5'HCV plasmid by replacing the

fragment between the XBa I site located upstream of the T7 promoter and the Nhe I site (nt 249-254) in the 5' NTR of HCV.

Cell cultures

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MDBK cells were obtained from M. Collett (ViroPharma, Inc.) and BT cells were obtained from the American Type Culture Collection (Rockville, Maryland). Cells were grown in Dulbecco's modified Eagle medium (D-MEM) supplemented with 10% horse serum and sodium pyruvate.

Transcriptions and transfections

All the designed plasmids, including pBVDV and the chimeric derivatives, were digested to completion with *Sda* I (*Sse*83871), purified by phenol extraction, precipitated by ethanol, and dissolved in water. The transcription reactions were performed sin the T7 Megascript kit (AMBION) using the conditions recommended by the manufacturer. Reactions were incubated at 37°C for 1 h, and ³H-UTP was added to the reaction to quantify the RNA synthesis. The quality of the synthesized RNAs was checked by agarose gel electrophoresis, and samples containing 50-60% of full-length RNA were used for electroporations and in vitro translations. The reaction mixtures were aliquoted and stored at -70°C prior to electroporation or in vitro translations.

Transfection was performed by electroporation of MDBK cells using previously described conditions (Mendez et al., 1998, supra). Two micrograms of in vitro synthesized RNA, corresponding to approximately 1 µ g of the full-length transcript, were used per electroporation. In standard experiments, ten-fold dilutions of electroporated cells were seeded in 6-well tissue culture plates containing 5 x 10⁵ naive MDBK cells per well. After 1 h of incubation at 37°C in an 5% CO₂ incubator, cells were overlaid with 3 ml of 0.6% LE Sea Kem agarose (FMC Bioproducts) containing minimal essential medium supplemented with 5% horse serum. Plaques were stained with crystal violet after 3 days incubation at 37°C. The rest of the transfected cells was seeded into 100-mm dishes and incubated for approximately 48 h or until cytopathic effect was observed in virtually all cells. Samples of the media were taken at 24 and 48 h, and virus titers were determined as described above and previously (Mendez et al., 1998, supra).

Analysis of the 5' ends of viral genomes

Sequencing of the 5' ends of selected variants of BVDV was performed on plaquepurified viruses. Plaques were typically isolated from the agarose overlay without staining with neutral red. Virus was eluted in 1 ml of D-MEM/10% horse serum for several hours and was used to infect 5 x 10⁵ MDBK cells in 35-mm dishes. After 1 h of virus adsorption of 37

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°C, an additional 1 ml of D-MEM/10% horse serum was added to the dishes, and incubation was continued for 36-48 h until cytopathic effect was observed in virtually all cells.

Fifty microliters of harvested viral stocks were clarified by low speed centrifugation, and viral RNAs were isolated by TRIzol reagent (Gibco-BRL) using the protocol recommended by the manufacturer. Sequencing of the 5' termini was performed using an oligonucleotide/cDNA-ligation strategy described elsewhere (Troutt et al., Proc. Natl. Acad. Sci. USA 89:9823-9825, 1992). The primer S1 (5'-GTCGTTTCACACATGGATCC), complementary to nt 710-729 of the BVDV genome, was used for cDNA synthesis. A phosphorylated oligonucleotide tag (5'-GACTGTTGTGGCCTGCAGGGCCGAATT) with an amino group on the 3' terminus was ligated to the first strand cDNA (Troutt et al., 1992, supra). One tenth of this reaction mixture was used for PCR amplification. The primers for PCR amplification were as follows: primer A (5'-GCCCTGCAGGCCACAACAGTC), complementary to the tag; primer B (5'-TCAGGCAGTACCACAA) complementary to nt 281-296 of the HCV 5' NTR; and primer C (5'-GGAATGCTCGTCAAGAAGACAG), complementary to nt 268-289 of the EMCV 5' NTR. The primer pairs of A + B or A + C were used for analysis of the pseudorevertants of 5'HCV and BVDV + HCVdelB1B2B3 or 5'EMCV, respectively. For the 5'HCV pseudorevertants, one tenth of the ligation mixture was used for an additional PCR reaction. This fragment was synthesized using primer S1, describe above, and a primer corresponding to nt 147-175 of the HCV genome. Fragments were purified by agarose gel electrophoresis and cloned into the plasmid pRS2. Multiple independent clones were sequenced by the standard dideoxy-mediated chain termination methods using the Sequenase version 2.0 DNA Sequencing Kit (USB).

Cell-free translation

Cell-free translation reactions were performed in reticulocyte extracts (Promega) using conditions recommended by the manufacture. Usually 0.1-1 μg of the same in vitro synthesized RNAs used in transfection experiments were used in 25 μl translation reactions. After 45 min of incubation at 30 °C, 2 μl were dissolved in 10 μl of sample buffer, and those samples were analyzed by sodium dodecyl sulfate PAGE. Labeled proteins were visualized by autoradiography of the dried gel. The efficiency of translation was measured using phosphorimager analysis (Molecular Dynamics) by comparing the radioactivity in the band corresponding to the N^{pro} protein. In preliminary experiments, an eightfold increase in incorporation was observed for translation of 4 μg versus 0.4 μg BVDV transcript RNA. Quantitative data were obtained from reactions using subsaturating (0.4 μg) amounts of BVDV or BVDV chimera transcript RNAs.

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Analysis of virus specific RNAs

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The protocols used for radioactive labeling of virus-specific RNAs are described in the appropriate figure legends. RNAs were isolated from the cells by using TRIzol reagent as recommended by the manufacturer (Gibco-BRL). After denaturation with glyoxal in dimethylsulfoxide, cellular RNAs were analyzed by electrophoresis in a 1% agarose gel containing a 10 mM phosphate buffer. Pieces of the dried gel containing the appropriate RNA bands were excised, and their radioactivity measured by liquid scintillation counting.

Results

Features of the BVDV, HCV, and EMCV 5' NTRs important for chimera design

Schematic representations of the proposed secondary structures of the 5' NTRs of HCV, BVDV, and EMCV are shown, and the location of each IRES is indicated in Figure 1. EMCV is a member of the cardiovirus genus within the family Picornaviridae. While not a member of the Flaviviridae, EMCV is similar to HCV and BVDV in that it is a positivestrand RNA virus shown to contain an IRES within its 5' NTR (Jang et al., J. virol 62:2636-2643, 1988). Based on their proposed secondary structures, the HCV IRES and the BVDV IRES have been classified as type 3 IRESs, while the EMCV IRES is classified as a type 2 IRES (Lemon & Honda, Siemin. Virol. 8:274-288, 1997). However, these three IRESs as well as IRESs from other members of the Flaviviridae and the Picornaviridae have been proposed to contain a common structural core (Le et al., Virus Genes 12:135-147, 1996).

The model for the secondary structure of the 341-nt HCV 5' NTR has been refined by enzymatic and chemical analysis of synthetic transcripts (Brown et al., Nucl. Acids. Res. 20:5041-5045, 1992; Wang et al., J. Virol 68:7301-7307, 1994; Honda et al., RNA 2:955-968, 1996; Lima et al., 1997). This element contains four discreet hairpins (referred to here as H1, H2, H3 and H4) and a pseudoknot at the base of hairpin H3 (Wang et al., 1995). The secondary structure of the 385-nt BVDV 5' NTR has not been as extensively studied, but is proposed to be similar to that of HCV (Brown et al., 1992) with four discrete hairpins (referred to here as B1', B1, B2, and B3) and a pseudoknot at the base of B3 (Rijnbrand et al., 1997). The secondary structure of the longer (>700 nt) EMCV 5' NTR consists of a series of hairpins A-M (Duke et al., 1992; Hoffman & Palmenberg, 1996). Recently, a revised model of the EMCV 5' NTR suggests moderately different secondary structures for the C and G subregions, and significantly different secondary structures for the I-M subregion (Palmenberg & Sgro, 1997).

For HCV, H1 is nonessential for IRES function (Reynolds et al., 1995; Rijnbrand et 35 al., 1995; Honda et al., 1996b; Reynolds et al., 1996; Kamoshita et al., 1997) and its deletion

has actually increased translation efficiency in some analyses (Rijnbrand et al., 1995; Honda et al., 1996b). Most studies have found that hairpin H2 and H3 and the pseudoknot are essential for IRES function (Wang et al., 1993; Rijnbrand et al., 1995; Honda et al., 1996b). However, two studies indicate that H2 may not be essential (Tsukiyama-Kohara et al., 1992; Urabe et al., 1997). The 3' boundary of the HCV IRES is more controversial. The IRES 5 clearly extends to the AUG initiation codon. However, some studies indicate that sequences affecting the efficiency of translation initiation extend into the ORF (Reynolds et al., 1995; Honda et al., 1996a; Honda et al., 1996b; Lu & Wimmer, 1996). By analogy to the HCV IRES and the related pestivirus CSFV IRES, the BVDV IRES probably requires hairpins B2 and B3 and the pseudoknot for function, with B1' and B1 probably not required for IRES 10 activity (Poole et al., 1995; Rijnbrand et al., 1997). For EMCV, hairpins H-L have been shown to be required for IRES function in mono- or dicistronic constructs (Jang & Wimmer, 1990; Duke et al., 1992). The remaining portion of the EMCV 5' NTR is thought to be required for RNA replication or unknown steps in viral replication that are important for pathogenesis (Duke et al., 1990; Martin & Palmenberg, 1996). 15

Replacement of the BVDV 5' NTR with the HCV 5' NTR results in a large decrease in specific infectivity

Since the BVDV 5' NTR and the HCV 5' NTR are proposed to have similar RNA secondary structure and functional organization, an experiment was performed to test whether 20 the BVDV 5' NTR could be replaced by the HCV 5' NTR. p5' HCV has an exact replacement of the BVDV 5' NTR with that of HCV (Fig. 2A) while the coding sequence and 3' NTR of p5'HCV are identical to pBVDV. Positioning of the HCV 5' NTR in such a manner was necessary since translation initiation from the HCV IRES begins at or near the AUG start codon (Honda et al., 1996a; Reynolds et al., 1995; Reynolds et al., 1996; Rijnbrand et al., 25 1996). The specific infectivity of 5'HCV RNA synthesized in vitro was compared to that of BVDV RNA by transfection of MDBK (bovine kidney) cells (Fig. 2A). The specific infectivity of BVDV RNA was approximately 4 x 106 plaque forming units (PFU)/µg RNA. In contrast, the specific infectivity of 5' HCV RNA was near the limit of detection (30-50 PFU/µg RNA) and considerable plaque heterogeneity was apparent. These results suggested 30 that the HCV 5' NTR replacement chimera might be incapable of efficient replication and plaque formation and that the plaque forming virus observed had arisen by secondary mutation(s). Sequence analysis of plaque-purified 5' HCV viruses presented below confirmed that the replicating pool of virus contained such pseudorevertants.

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Next, the *in vitro* translation efficiency of these two RNAs in rabbit reticulocyte extracts was analyzed to test whether the defect in specific infectivity of 5' HCV RNA could be attributed to lower translation efficiency. Although the specific infectivity of 5' HCV RNA was reduced ~5 logs compared to BVDV RNA, its translation efficiency was only slightly reduced, ~twofold (Fig. 3, lane 1 vs. lane 2). The apparent size of the N-terminal cleavage product, N^{pro}, was identical for both RNAs, suggesting that translation initiated with the correct AUG. These data are consistent with the hypothesis that the BVDV 5' NTR contains signals that are required for a step in replication other than translation which are not present in the 5' HCV chimera.

Given the low specific infectivity of 5' HCV RNA, an experiment was performed to test the effect of placing the BVDV 5' NTR sequence upstream of the HCV 5' NTR, resulting in tandem BVDV and HCV 5' NTRs (called BVDV + HCV). This arrangement actually decreased translation efficiency (Fig. 3, lane 14 vs. lane 1) yet restored infectivity (Fig. 2A). The plaques produced by BVDV + HCV were also heterogeneous in size, indicating that this virus was unstable. Upon passage, RT-PCR analysis indicated that pseudorevertants had indeed arisen in which portions of the BVDV and/or HCV 5' NTRs had been deleted (data not shown). These data show that sequences in the BVDV 5' NTR required for virus replication can function when placed upstream of a functional HCV IRES driving translation of the BVDV polyprotein.

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Hairpins B1' and B1 in conjunction with the HCV IRES are sufficient for stable and efficient BVDV replication

The sequences within the BVDV 5' NTR that restored replication in the context of the HCV 5' NTR were mapped using three deletion variants. The deletion BVDV + HCVdelB3 removed a large portion of hairpin B3; the deletion within BVDV + HCVdelB2B3 removed hairpins B2 and B3, and the deletion within BVDV + HCVdelB1B2B3 removed hairpins B1, B2 and B3. The specific infectivities of RNAs from these deletion mutants were near that of BVDV RNA (Fig. 2). Upon passage of these viruses, RT-PCR analyses and sequencing indicated that BVDV + HCV delB3 and BVDV + HCVdelB2B3 were stably propagated and produced homogeneous plaques slightly smaller than those of wild-type BVDV (data not shown). In contrast, BVDV + HCVdelB1B2B3 produced smaller heterogeneous plaques. Reverse transcription-polymerase chain reaction (RT-PCR) analysis and sequencing indicated that BVDV + HCVdelB1B2B3 underwent a reversion event described in more detail below. The translation efficiencies of these three RNAs (Fig. 3, lanes 9, 10, and 12) were similar to BVDV + HCV RNA (Fig. 3, lane 14), indicating that the deleted portions (hairpins B1, B2,

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and B3) are not required for translation in the BVDV + HCV chimera. These results show that B1' and B1 are the minimal elements sufficient for stable replication in conjunction with the HCV 5' NTR.

Having shown that B1' and B1 are sufficient for replication in conjunction with the HCV 5' NTR, we next conducted a deletion analysis to determine the sequences within the HCV 5' NTR of BVDV + HCV delB2B3 required for replication. A large portion of H1 was deleted in BVDV + HCV delB2B3H1, while both H1 and H2 were deleted in BVDV + HCV delB2B3H1H2. Of these two RNAs, only BVDV + HCV delB2B3H1 was as infectious as parental BVDV RNA (Fig. 2B). However, the BVDV + HCV delB2B3H1 virus produced smaller plaques than BVDV + HCV delB2B3, indicating that hairpin H1 may augment replication of the chimera. In contrast, BVDV + HCV delB2B3H1H2 RNA was not infectious (Fig. 2B) and was translated poorly (Fig. 3, lane 11). Diminished HCV IRES activity might be due to deletion of hairpin H2 or juxtaposition of BVDV hairpins B1' and B1 with H3. A third derivative of BVDV + HCV delB2B3, with a Sma I-Sma I deletion abrogating HCV IRES function by removing H3, was also not infectious (data not shown). Thus, a 5' NTR consisting of B1' and B1 and a functional HCV IRES is sufficient for stable BVDV replication in MDBK cells. Similar results were obtained in BT cells, another BVDV-permissive continuous bovine cell line (data not shown).

20 Replacement of the BVDV 5' NTR with the EMCV 5' NTR

The following experiment was performed to determine whether the BVDV 5' NTR could be replaced by the 5' NTR of a more phylogenetically distant virus, EMCV. A derivative of BVDV was created, called 5' EMCV, that contains an exact replacement of the BVDV 5' NTR with the EMCV 5' NTR plus an additional guanosine residue at the 5' terminus for more efficient transcription initiation of T7 polymerase (Fig. 4A). The specific infectivity of 5' EMCV RNA was more than three orders of magnitude lower than BVDV RNA, indicating that it was defective for replication, although its specific infectivity was higher than that of 5' HCV RNA (compare Figs. 4A and 2A). Similar to 5' HCV, 5' EMCV produced heterogeneous plaques, and sequence analysis indicated that pseudorevertants had arisen. The lower specific infectivity of 5' EMCV RNA was not likely because of a defect in translation, since the translation efficiency of 5' EMCV RNA was about threefold higher in vitro than that of BVDV RNA (Fig. 3, lane 20 vs. lane 19).

Similar to BVDV + HCV, it was also determined whether the BVDV 5' NTR at the 5' end of the 5' EMCV RNA would increase its specific infectivity. BVDV + EMCVdelA (Fig. 4A) contained the entire BVDV 5' NTR in tandem with the EMCV 5' NTR lacking a portion

of hairpin A. BVDV + EMCVdelA RNA had a specific infectivity near that of BDVD RNA (compare Figs. 4A and 2A) despite having a lower translation efficiency than 5' EMCV (Fig. 3, lane 21 vs. lane 20). Similar to the results with BVDV + HCV, this implicates the added BVDV 5' NTR sequence for a step in viral replication other than translation. Two derivatives of BVDV + EMCVdelA that contain deletions of portions of the BDVD 5' NTR but maintain the sequence of B1' and B1, BDVD + EMCVdelB3A and BVDV + EMCVdelB2B3A (Fig. 4A), also were infectious. These derivatives had translation efficiencies near that of the parental BVDV + EMCVdelA (Fig. 3, compare lanes 15 and 16 with lane 21). This demonstrated that hairpins B1' and B1 were sufficient for replication in conjunction with a large portion of the EMCV 5' NTR. Derivatives of BVDV + EMCVdelB3A or BVDV + EMCVdelB2B3A that contain further deletions of EMCV (BVDV EMCVdelB3ABC and BVDV + EMCVdelB2B3ABC in particular) were translated efficiently (Fig. 3, lanes 17 and 18) and were infectious (Fig. 4B). This indicates that the chimeras did not require putative EMCV RNA replication signals (Martin & Palmenberg, 1996). However, derivatives with deletions extending into the canonical EMCV IRES were not infectious. For example, BVDV + EMCVdelB3A-H and BVDV + EMCVdelB2B3A-H, in which a portion of hairpin H is deleted, were not infectious (Fig. 4B) and were inefficiently translated in vitro (Fig. 3, lanes 22 and 23). It should be noted that all of the BVDV + EMCV chimeras produced plaques of heterogeneous size, indicating some instability.

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Relatively simple 5' NTR mutations are observed in adapted pseudorevertants

As mentioned previously, BVDV + HCVdelB1B2B3 did not replicate stably as indicated by the heterogeneity in the size of plaques produced by this virus. Upon passage and selection of medium plaque-producing variants, 5' RACE analysis and sequencing indicated that nt 1-26 had been deleted in the pseudorevertants, removing a large portion of B1' which was apparently deleterious in the absence of B1. This deletion results in the 5' terminal sequence 5'GUAUCG which is identical to the first six bases of BVDV genome RNA (Fig. 5) and is repeated at positions 27-32.

Analysis of the passaged 5' EMCV virus indicated that the replicating progeny had also undergone a simple deletion of sequence at the 5' end to generate more efficiently replicating variants (Fig. 5). After electroporation, the 5' EMCV virus pool was passaged 5 times at a multiplicity of infection of 0.1-1 PFU/cell on MDBK or BT cells, and the 5' termini of three randomly picked plaques were sequenced. For all three plaques selected, nt 2-209 had been deleted, again creating a genome RNA with the 5' terminal tetranucleotide sequence 5'-GUAU.

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Analysis of the 5' HCV progeny indicated that more complicated variants had arisen. Most small plaque-producing variants were unstable and quickly reverted to medium plaqueproducing variants. However, one small plaque-producing variant and two stable medium plaque-producing variants were isolated. 5' terminal sequences of the variants were amplified by rapid amplification of cDNA ends (RACE) and cloned into a plasmid vector, and sequences for several independent colonies were determined. The sequence of three clones of the small plaque-producing virus (5'HCV.R1) contained a deletion of HCV sequence from nt 1-34 and an addition of the dinucleotides 5'-AU in two clones and 5'-GU in the third clone. This creates a 5' terminus of 5'-(G/A) UAA (Fig. 5B), reminiscent of the first three bases of the BVDV genome RNA (5'-GUA). Both medium plaque variants appeared to have arisen by RNA recombination with non-viral sequences (Fig. 5). One medium plaque variant (5' HCV.R2) had deleted the first 21 bases of the HCV sequence and contained instead a heterologous sequence of 22 bases. BLAST searches revealed a perfect match between this sequence and a sequence in a human retina cDNA of unknown function (Tsp509I). The second medium plaque variant (5' HCV.R3) had also undergone a possible recombination event leading to the addition of 12 nt to the 5' end of the HCV sequence. Given its short length, multiple matches were found in the database with this sequence. As for the small plaque variant, sequencing of multiple clones revealed heterogeneity out the extreme 5' end. with either G of A identified as the 5' base. Remarkably, for both medium plaque variants, the fused heterologous sequence began with the tetranucelotide sequence 5'-(G/A) UAU (Fig. 5B). For all three variants, sequencing of the entire 5' NTR and a portion of the N^{pro} coding region revealed only these changes at the 5' termini.

5' NTR sequence changes are sufficient for the pseudorevertant phenotypes

To assess the importance of these alterations oat the 5' terminus of the 5' HCV pseudorevertants, derivatives of 5' HCV were created with the changes determined by 5' RACE (Fig. 6A) and analyzed the specific infectivities of these RNAs (Fig. 6B). Corresponding to the small plaque variant, a derivative called 5' HCV.R1 orig was engineered which contained a 5' NTR consisting of the dinucleotide 5' -GU at the 5' terminus of HCV nt 35-341. This results in a 5' terminus consisting of 5'-GUAA. 5'HCV.R1 orig RNA had a specific infectivity at least four orders of magnitude higher than 5' HCV RNA (Figs. 6B and 2A). This demonstrates that this 5' NTR structure is sufficient for phenotypic reversion to high specific infectivity. However, small plaques and considerable heterogeneity were observed for 5'HCV.R1 orig suggesting that additional mutations may be present in the original small plaque variant.

The engineered derivative 5'HCV.R2orig had a 5' NTR consisting of 22 nt of Tsp509I-homologous sequence followed by HCV nt 22-341. Another construct, called 5'HCV.R3orig was made, which has the 12 nt of the other heterologous sequence fused to the intact HCV 5' NTR. Specific infectivities for both these derivatives were essentially the same as observed for wild type BVDV RNA (2-4 x 10⁶ PFU/µg; Fig. 6B). Transfection with these transcripts produced medium plaques, as observed for the original variants, and this phenotype was stable upon passaging. These results show that the altered 5'NTR sequences were responsible for the pseudorevertant phenotypes rather than changes elsewhere in their genomes.

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Addition of the tetranucleotide sequence 5'-GUAU to the HCV 5' NTR allows efficient BVDV replication

For all three 5' HCV variants studied, as well as the BVDV + HCV delB1B2B3 and 5'EMCV pseudorevertants, 5' NTR alterations seemed to involve creation of a three- or fourbase "consensus" sequence identical to the 5' terminus of BVDV genome RNA. To test the importance of this sequence, as opposed to fused heterologous sequences, we created a set of variants with the BVDV 5' tetranucleotide sequence linked to the HCV 5' NTR or the deletion/recombinant break points identified during sequence analysis of the 5' HCV pseudorevertants (Fig. 6A). 5' HCV.R1cons had the tetranucleotide sequence 5'-GUAU fused to HCV nt 35-341. 5'HCV.R2cons had the 5'-GUAU tetranucleotide sequence fused to HCV nt 22-341. 5'HCV.R3cons contained the tetranucleotide sequence 5'-Guau fused to the intact 5' terminus of the HCV NTR. RNAs from all three of these derivatives had specific infectivities more than five orders of magnitude higher than 5'HCV and comparable to parental BVDV (Fig. 6B).

There were, however, significant differences between the phenotypes of some of these derivatives versus the reconstructed pseudorevertants. As mentioned above, 5'HCV.R1orig yielded tiny and small plaques and produced low virus yields even after 48 h. In contrast, the addition of four bases rather than two bases (5'-GUAU vs. 5'-GU) yielded virus with near wild-type plaque morphology (Fig. 6B) and growth Rates (Fig. 7). In the case of the smaller deletion, 5'HCV.R2orig and 5'HCV.R2cons were indistinguishable, suggesting that, other than the 5' four bases, the fused heterologous sequences were dispensable. This was not he case, however, for the chimera containing the 5'-GUAU tetranucleotide sequence

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fused to the intact HCV 5' NTR. 5'HCV.R3cons produced small plaques (Fig. 6B) and grew more slowly than 5'HCV.R3orig (Fig. 7) suggesting that the sequence/structure of the sequences downstream of the 5' four bases can affect replication efficiency.

5 The tetranucleotide sequence 5'-GUAU is important for efficient BVDV RNA accumulation

Next, the effects of the different 5' termini on virus-specific RNA accumulation directly after transfection were analyzed. This allowed a direct comparison between 5'HCV and the reconstructed pseudorevertants as well as selected BVDV + HCV deletion constructs. MDBK cells were transfected with in vitro synthesized RNAs and labeled for 10 h beginning at 5 h post-transfection with ³H-UTP in the presence of actinomycin D (Fig. 8). RNA replication of the 5' HCV chimera was severely impaired to a level below detection (Fig. 8, lane 2). In contrast, every 5' NTR alteration of 5' HCV that increased RNA specific infectivity and allowed efficient virus growth led to readily detectable viral RNA accumulation. Addition of B1' and B1 to the 5' terminus of the HCV 5' NTR restored RNA replication to a level ~50% of that observed for BVDV (BVDV + HCVdelB2B3; Fig. 8, lane 3 vs. lane 1). BVDV + HCVdelB2B3H1 displayed reduced RNA synthesis compared to BVDV + HCVdelB2B3 (Fig. 8, lane 4 vs. lane 3) perhaps explaining its small plaque phenotype and suggesting a possible positive role for H1 in replication of this chimera. 5'HCV.R1orig, which had exhibited plaque heterogeneity and slow growth, accumulated less RNA when compared to 5'HCV.R1cons (Fig. 8, lane 5 vs. lane 6). 5'HCV.R2orig and 5'HCV.R2cons showed similar RNA accumulation (Fig. 8, lane 9 vs. lane 10) consistent with their medium plaque phenotypes; and 5HCV.R3cons exhibited reduced RNA synthesis compared to 5'HCV.R3orig (Fig. 8, lane 8 vs. lane 7), consistent with their small-versus medium-plaque phenotypes.

Although these RNA phenotypes are complex, the most striking result is that addition of the B1' B1 hairpins, addition of heterologous 5' sequences terminating with 5'-GUAU or simply fusion of this tetranucleotide sequence with the HCV 5' NTR or short 5' truncations of the HCV 5' NTR all dramatically upregulated RNA accumulation. This occurred without increasing translation efficiency, at least as measured in a cell-free assay (Fig. 3, compare lanes 3-8 to lane 1), suggesting that these sequences function at the level of RNA replication or stability.

Discussion

The work presented here helps to define the requirements for a functional BVDV 5'NTR. The BVDV-specific 5' NTR sequences required for efficient replication in cell culture are minimal and consist of the 5' terminal sequence, 5'-GUAU. The sequence 5'-AUAU, detected for some pseudorevertants, may also be functional but this was not tested for technical reasons. This simple 5'-terminal tetranucleotide sequence, which is conserved among pestivirses (Ruggli et al., 1996; Becher et al., 1998), was shown to function in the context of functional IRES elements derived from the hepacivirus HCV or the picornavirus EMCV. As discussed below, this may indicate that the 5' signals required for BVDV RNA replication are rather simple or that elements in these heterologous IRESs can functionally replace deleted BVDV sequences.

Sequences at the extreme 5' end of BVDV genome RNA could modulate the efficiency of RNA accumulation by affecting RNA stability, translation, promoter efficiency, or some combination of these processes. At this time, we can not distinguish among these possibilities but favor an effect on RNA replication. The complement of the BVDV 5' sequence at the 3' end of the negative-strand RNA presumably functions in the initiation of positive-strand RNA synthesis. Thus, AUAC-3' at the 3'terminus fo minus-strand RNA may be important for positive-strand RNA synthesis. Interestingly, for some positive-strand RNA viruses such as rubella virus (Pugachev & Frey, 1998), flock house virus (Ball, 1994) and turnip crinkle virus (Guan et al., 1997), only minimal cis-acting sequences at the 3' termini of negative-strand RNAs are required positive-strand RNA synthesis. In contrast to the 5' NTR replacements, we were unable to generate replication-competent BVDV-HCV replacing that of BVDV (data not shown). This may indicate that the signals within the pestivirus 3' NTR required for initiation of negative-strand RNA synthesis are more complex and virus specific. Once the replication complex has assembled at the 3' NTR and transversed the RNA during negative-strand synthesis, the requirements of the 5' NTR for initiation of positive-strand synthesis may be minimal.

Although the RNA replication signals within the 5' NTR appear to be rather simple, it is possible that the signals important for RNA replication actually extend into the IRES and are more complicated. For instance, the 5'HCV pseudorevertants were more stable and grew to higher titers than the 5'EMCV counterparts, despite the fact that the 5'EMCV RNAs were translated more efficiently in vitro. This may indicate that the BVDV and HCV IRESs contain signals important for RNA synthesis that are absent in the EMCV IRES.

It is perhaps not surprising that 5' HCV appeared to recombine with cellular mRNAs to acquire a 5' terminus with the 5' -(G/A) UAU consensus, given that non-cytopathic strains

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of BVDV can recombine with BVDV RNA or cellular mRNAs to generate cytopathic strains of BVDV (Meyers & Thiel, 1996). Presumably, this recombination event involves template switching during negative-strand RNA synthesis, as observed for polio-virus (Kirkegaard & Baltimore, 1986). In contrast to 5' HCV, simple deletions of 5' terminal viral sequences could account for the BVDV + HCVdelB1B2B3 and 5'EMCV pseudorevertants since the 5 tetranucleotide sequence is present in these 5' NTRs upstream of functional IRES elements. Such deletions could occur by partial degradation of positive-strand template prior to negative-strand synthesis, by premature termination during negative-strand RNA synthesis, or by degradation of 3' terminal negative-strand sequence after synthesis. It is proposed that 5'HCV was forced to recombine with cellular sequences because HCV does not have an 5'-10 (G/A) UAU sequence upstream of its IRES. The first occurrence of an (G/A)UAUA tetranucleotide sequence is at nt 94-97 within hairpin H2, and a 5' deletion extending into this sequence would presumably inactivate or severely impair HCV IRES activity. It is interesting that BVDV + HCVdelB1B2B3 and 5'EMCV pseudorevertants were generated at much higher frequency than 5'HCV pseudorevertants. This may indicate that recombination between 15 BVDV and cellular RNAs is a rare event compared to the processes which lead to deletion of terminal viral sequences.

Poliovirus chimeras dependent upon a functional HCV IRES have been reported (Lu & Wimmer, 1996). Interestingly, viable poliovirus chimeras were produced only when HCV sequences included both the IRES and the N-terminal portion of the HCV ORF. Nucleotide 20 sequences or structures in the downstream ORF can modulate HCV IRES translational efficiency (see Reynolds et al., 1995; Honda et al., 1996a) but it was also suggested that the N-terminal portion of the HCV core polypeptide might be involved. In the case of our 5' HCV pseudorevertants, there is no requirement for HCV C protein sequences. Although the 25 translation efficiency of the HCV IRES in the presence of additional HCV sequences 3' to the AUG start was not directly assessed, the HCV chimeras and pseudorevertants were translationally active and infectious in the absence of any portion of the HCV ORF. This indicates that either the HCV IRES does not extend into the HCV ORF or that the BVDV ORF contains analogous sequence which functions in our 5HCV chimeras. There is some limited identity between HCV and BVDV within this region. For example, HCV nt 359-394 30 and BVDV nt 405-440 are identical at 21 of 36 positions, although identity within this sequence may be attributed to a high adenosine content. It is interesting to note that the luciferase (LUC) and chloramphenicol acetyl transferase (CAT) reporter genes previously used to detect HCV IRES activity (Tsukiyama-Kohara et al., 1992; Wang et al., 1993) also 35 have adenosine- or purine-rich regions in relatively the same position as the HCV ORF and

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BVDV ORF. It this region is indeed important for IRES activity, this may explain why some have observed that the HCV IRES does not require a portion of the HCV ORF for translation of CAT or LUC (Tsukiyama-Kohara et al., 1992; Wang et al., 1993). Point mutations and insertions within this region of HCV have been shown to reduce HCV IRES activity in vitro (Honda et al., 1996a,b).

Despite the fact that B1' and B1 are conserved among different strains of BVDV and similar hairpins are present in border disease virus and CSFV (Deng & Brock, 1993; Becher et al., 1998), B1' and B1 were dispensable for BVDV replication, provided that the 5' tetranucleotide sequence 5'-(G/A)UAU remained. This may indicate a role for B1' and B1 in viral replication in vivo that we do not observe in cell culture. It will be interesting to test the phenotype of chimeras that lack B1' and B1 in vivo to determine if they are attenuated and might serve as useful BVDV vaccines. In this vein, several studies with flaviviruses have demonstrated that alterations in 5' NTR or 3' NTR elements can lead to attenuation in vivo (Cahour et al., 1995; Men et a., 1996; Mandl et al., 1998). BVDV chimeras that utilize the HCV or EMCV IRES may also prove to be attenuated simply due to the presence of the heterologous IRES. For poliovirus, it has been shown that differences in IRES efficiency in different host-cell environments can modulate host range and virulence (Shiroki et al., 1997).

BVDV-HCV chimeras that are dependent on a functional HCV IRES may have another practical application. It may be possible to use these chimeras to screen for anti-HCV therapeutics that target the HCV IRES. Other researchers have shown antisense oligonucleotide-mediated inhibition of HCV gene expression in hepatocytes by targeting the oligonucleotides to the HCV IRES (Hanecak et al., 1996). It will be of interest to measure the efficacy of antisense oligonucleotides or ribozymes (Lieber et al., 1996) against replicating virus, and these chimeras are more useful than HCV for this purpose since they are able to replicate efficiently in cell culture. BVDV is believed to be a reasonable model of HCV replication not only because of homology and conserved motifs within the 5' NTR but also because of similarities in overall genetic organization (Rice, 1996) and polyprotein processing strategy (Tautz et al., 1997; Xu et al., 1997).

In view of the above, it will be seen that the several advantages of the invention are achieved and other advantageous results attained.

As various changes could be made in the above methods and compositions without departing from the scope of the invention, it is intended that all matter contained in the above description and shown in the accompanying drawings shall be interpreted as illustrative and not in a limiting sense.

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All references cited in this specification, including patents and patent applications, are hereby incorporated by reference. The discussion of references herein is intended merely to summarize the assertions made by their authors and no admission is made that any reference constitutes prior art. Applicants reserve the right to challenge the accuracy and pertinency of the cited references.

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What is Claimed is:

- 1. A polynucleotide comprising a chimeric viral RNA which comprises:
- (a) a 5' nontranslated region (5' NTR);
- (b) an open reading frame (ORF) region; and
- 5 (c) a 3' nontranslated region (3' NTR);

wherein at least one of said regions is chimeric and comprises a first nucleotide sequence from a pestivirus in operable linkage with a first nucleotide sequence from an hepatitis C virus (HCV), and wherein said chimeric viral RNA is replication-competent.

- 10 2. The polynucleotide of claim 1, wherein the chimeric region is the 5' NTR and the first pestivirus nucleotide sequence is from a bovine viral diarrhea virus (BVDV).
 - 3. The polynucleotide of claim 2, wherein the BVDV nucleotide sequence is located at the 5' terminus of the chimeric 5' NTR and comprises 5' RUAU.

4. The polynucleotide of claim 3, wherein the first HCV nucleotide sequence in the chimeric 5' NTR comprises an internal ribosome entry site (IRES).

- 5. The polynucleotide of claim 4, wherein the ORF and the 3' NTR consist of second and third BVDV sequences.
 - 6. The polynucleotide of claim 5, wherein the 5' terminal sequence comprises 5' GUAU.
- 7. The polynucleotide of claim 4, wherein the ORF comprises a second HCV sequence encoding at least one structural protein operably linked to a second BVDV sequence.
- 8. The polynucleotide of claim 1, wherein the pestivirus is BVDV and the 30 chimeric region is the 3' NTR.
 - 9. The polynucleotide of claim 8, wherein the first HCV sequence in the chimeric 3' NTR comprises the HCV 98 bp 3' terminal element (SEQ ID NO:X) operably linked to the first BVDV sequence.

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- 10. A method for identifying compounds having antiviral activity against hepatitis C virus (HCV) comprising the steps of:
- (a) providing a first cell containing a chimeric viral RNA which is replication-competent in the cell, the chimeric viral nucleic acid comprising a 5' nontranslated region (5' NTR), an open reading frame (ORF) region; and a 3' nontranslated region (3' NTR); wherein at least one of said regions is chimeric and comprises a first nucleotide sequence from a pestivirus in operable linkage with a first nucleotide sequence from an hepatitis C virus (HCV);
 - (b) providing a second cell containing the pestivirus; and
- (c) comparing the replication efficiency of the chimeric viral RNA acid in the presence and absence of a test compound to the replication efficiency of the pestivirus in the presence and absence of the test compound, wherein a greater reduction in compound-induced replication efficiency of the chimeric viral RNA than the pestivirus indicates the compound has anti-HCV activity.

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- 11. The method of claim 10, wherein the chimeric region is the 5' NTR and the first pestivirus nucleotide sequence is from a bovine viral diarrhea virus (BVDV).
- 12. The method of claim 11, wherein the BVDV nucleotide sequence is located 20 at the 5' terminus of the chimeric 5' NTR and comprises 5' RUAU.
 - 13. The method of claim 12, wherein the first HCV nucleotide sequence in the chimeric 5' NTR comprises an internal ribosome entry site (IRES).
- 25 14. The method of claim 13, wherein the ORF and the 3' NTR comprise second and third sequences from the BVDV.
 - 15. The method of claim 10, wherein the pestivirus is BVDV and the chimeric region is the 3' NTR.

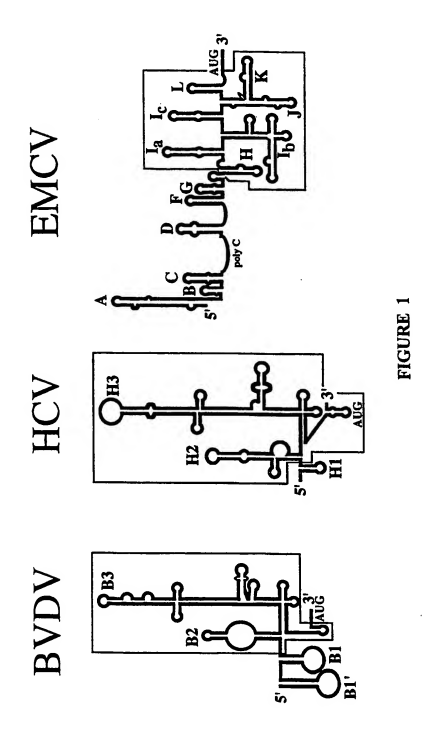
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- 16. A genetically-engineered virus comprising a chimeric RNA genome which comprises:
 - (a) a 5' nontranslated region (5' NTR);
 - (b) an open reading frame (ORF) region; and
- 35 (c) a 3' nontranslated region (3' NTR);

wherein at least one of said regions is chimeric and comprises a first nucleotide sequence from a pestivirus in operable linkage with a first nucleotide sequence from an hepatitis C virus (HCV), and wherein said chimeric RNA genome is replication-competent.

- 5 17. The genetically-engineered virus of claim 16, wherein the chimeric region is the 5' NTR and the first pestivirus nucleotide sequence is from a bovine viral diarrhea virus (BVDV).
- 18. The genetically-engineered virus of claim 16, wherein the BVDV nucleotide sequence is located at the 5' terminus of the chimeric 5' NTR and comprises 5' RUAU and the first HCV nucleotide sequence in the chimeric 5' NTR comprises an internal ribosome entry site (IRES).
- 19. A vaccine against bovine viral diarrhea virus (BVDV) comprising an
 15 immunogenically-effective amount of a genetically-engineered virus comprising a chimeric RNA genome having:
 - (a) a 5' nontranslated region (5' NTR);
 - (b) an open reading frame (ORF) region; and
 - (c) a 3' nontranslated region (3' NTR);
- wherein at least one of said regions is chimeric and comprises a first nucleotide sequence from BVDV in operable linkage with a first nucleotide sequence from an hepatitis C virus (HCV), and wherein the genetically-engineered virus is attenuated as compared to BVDV.
- 20. The vaccine of claim 19, wherein the chimeric region is the 5' NTR and the BVDV nucleotide sequence is located at the 5' terminus of the chimeric 5' NTR and comprises 5' RUAU and the first HCV nucleotide sequence in the chimeric 5' NTR comprises an internal ribosome entry site (IRES).
 - 21. A polynucleotide comprising a chimeric viral RNA which comprises:
- 30 (a) a 5' nontranslated region (5' NTR);
 - (b) an open reading frame (ORF) region; and
 - (c) a 3' nontranslated region (3' NTR);

wherein at least one of said regions is chimeric and comprises a first nucleotide sequence from a pestivirus in operable linkage with a heterologous nucleotide sequence and wherein said chimeric viral RNA is replication-competent.



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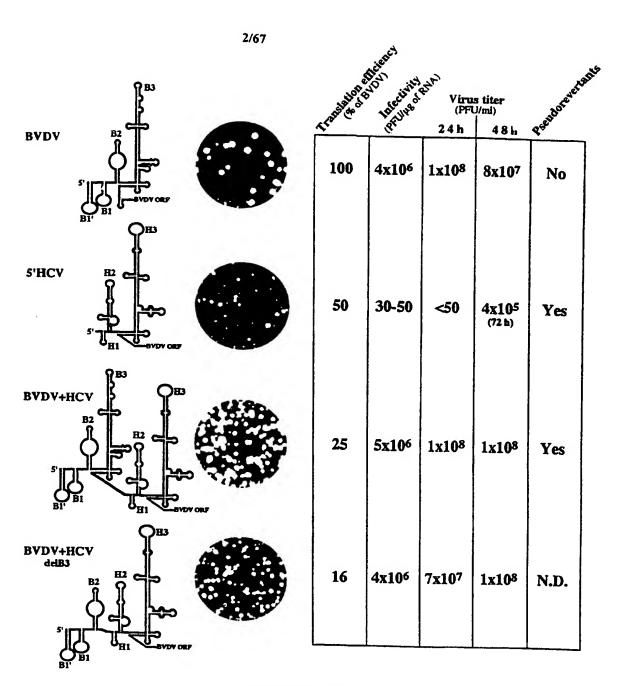


FIGURE 2A

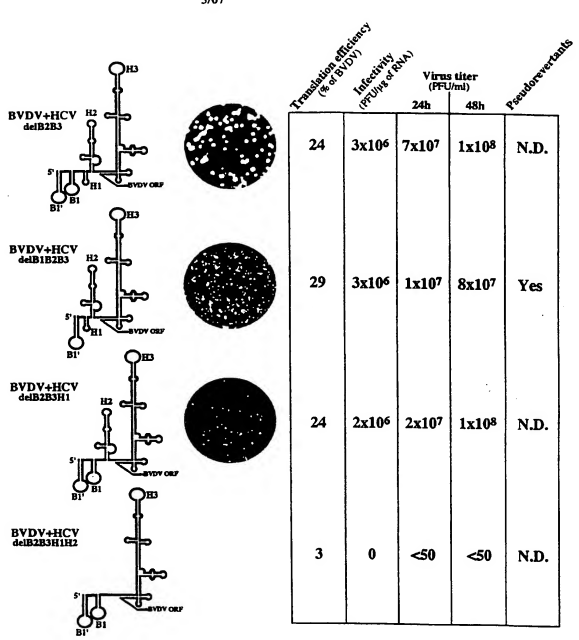
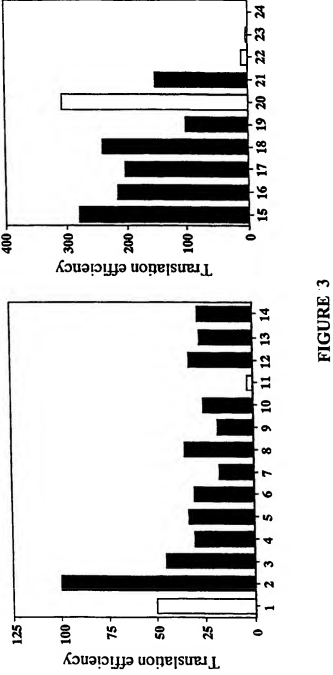


FIGURE 2B



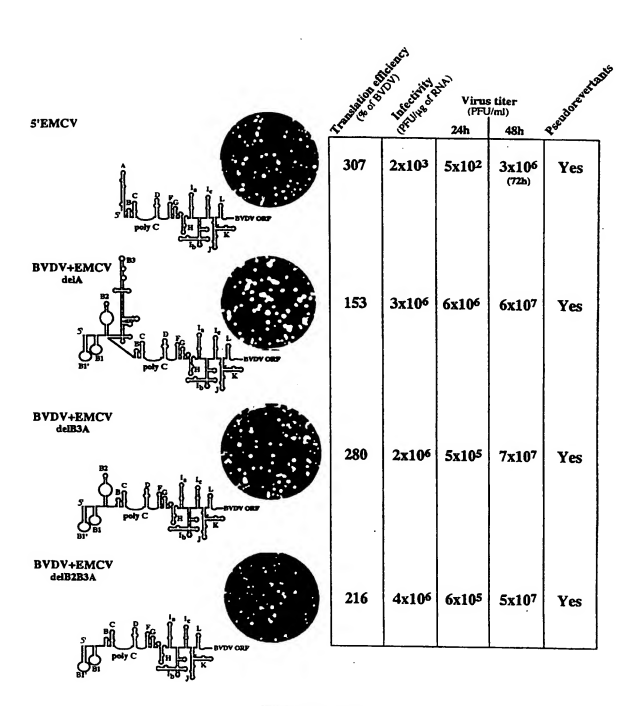


FIGURE 4A

menonin, asin meranca +

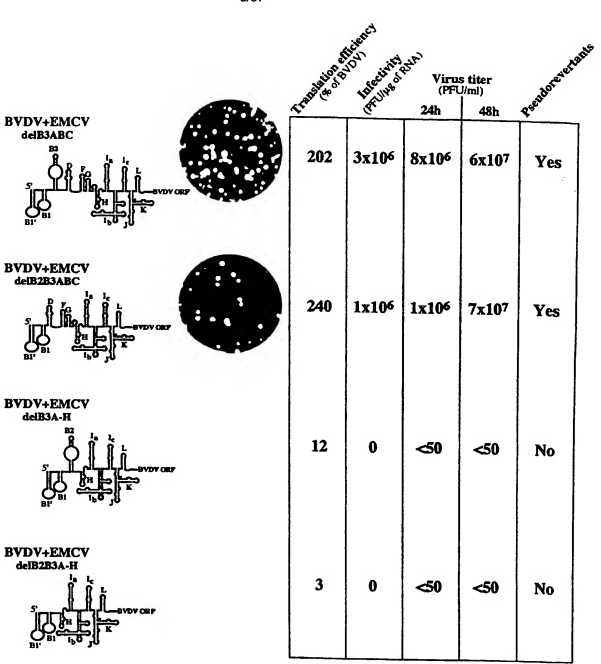
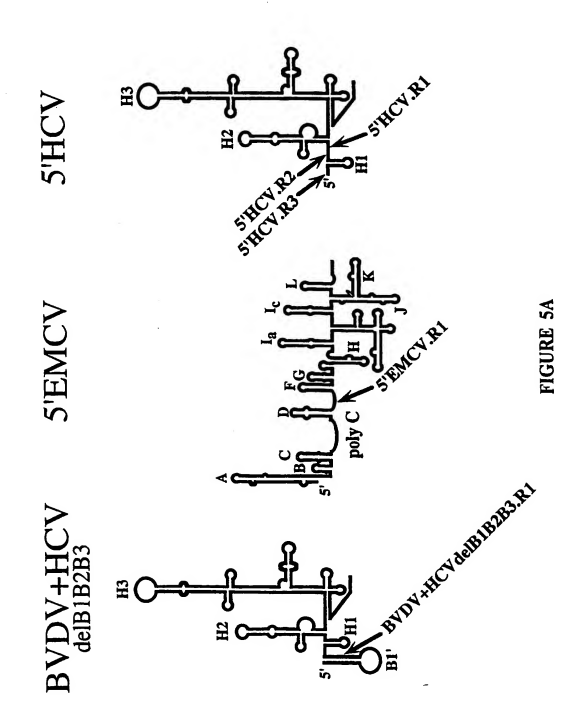


FIGURE 4B



UNICHOSTIC MAIO COCCACO I

guauacgagaauuagaaaaggcacucguauacguaCAUGGCACGUgccagcccccugauggggggc guauacguaCAUGGCACGUgccagcccccugaugggggc B1 . BVDV+HCVdelB1B2B3.R1 BVDV+HCVdelB1B2B3

BVDV+HCVdelB1B2B3

(G/A) Taaucacuccccugugaggaac gecagececeugauggggggggeacaeuceceaugaaucaeuceceugugaggaae Ħ 5'HCV.R1 5'HCV 5 · HCV

(G/A) UAUCAGAAGUGCGAAUGCUGAacacuccaccaugaaucacuccccugugaggaac (G/A) **UNTUGCAGUTU**gccagcccccugauggggggggcgacacuccaccaugaaucacuccccugugaggaac 5' HCV. R2 5' HCV. R3

S'EMCV

Guaugunauuuccaccauauug · · guungucuauauguuauuuuccaccauauug gungaaagccgggggggggggc..... 5'EMCV.R1 5'EMCV

FIGURE 5B

gccagcccccugauggggggggcacaccaccaugaaucacuccccugugaggaacu

GUaaucacuccccugugaggaacu

GUAUaaucacucccugugaggaacu

GUAUCAGAAGGCGAAUGCUGAacacuccaccaugaaucacuccccugugaggaacu

GUAU acacuccaccaugaaucacuccccugugaggaacu **<u>euau</u>ugcaeuu**gccagccccugauggggggggggcgacauccaccaugaaucacucccugugaggaacu

GUALgccagccccugaugggggggggacacuccaccaugaaucacuccccugugaggaacu

FIGURE 6A

5'HCV. Rlorig

5 HCV

5'HCV. R1cons

5 HCV. R2orig

5'HCV. R2cons

5'HCV. R3orig

5'HCV. R3cons

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	1 (a. <u>1</u>)	Translation	Infectivity	Virus tite	er (PFU/ml) j
	CAU.	efficiency (% of BVDV)	(PFU/µg of RNA)	24b	48h
BVDV		100	4x10 ⁶	7x10 ⁷	1x10 ⁸
5'HCV.R1orig		45	4x10 ⁵	2x10 ³	2x10 ⁵
5'HCV.R1cons		29	3x106	4x10 ⁷	5x10 ⁷
5'HCV.R2orig		17	2x10 ⁶	7x10 ⁶	5x10 ⁷
5'HCV.R2cons		35	3x10 ⁶	2x10 ⁷	4x10 ⁷
5'HCV.R3orig		33	3x10 ⁶	4x10 ⁷	5x10 ⁷
5'HCV.R3cons		30	3x10 ⁶	1x10 ⁷	6x10 ⁷

FIGURE 6B

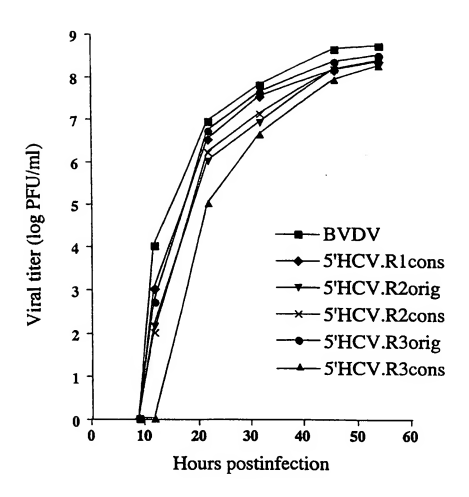
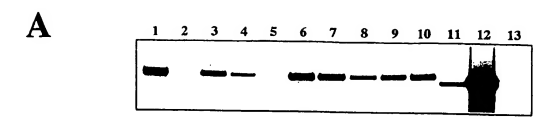


FIGURE 7



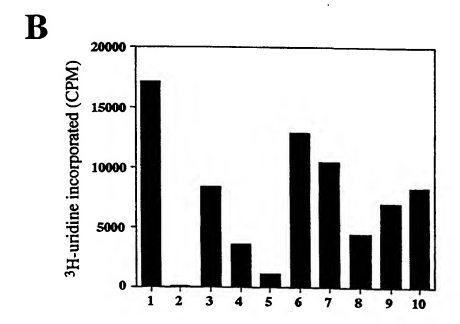


FIGURE 8

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pACNR/BVD NADL-Xba* -> Graphic Map

DNA sequence 15065 bp gtatacgagaat ... cgactcactata circular

packnr/BVD NADL-Xba = HaeII and XhoI digest of packnr/BVD NADL ligated to HaeII and XhoI digest of packnr1180/DraIII-/BVD5* 8/27 corrected nt 12136 G to C to give HpaI site.

Co

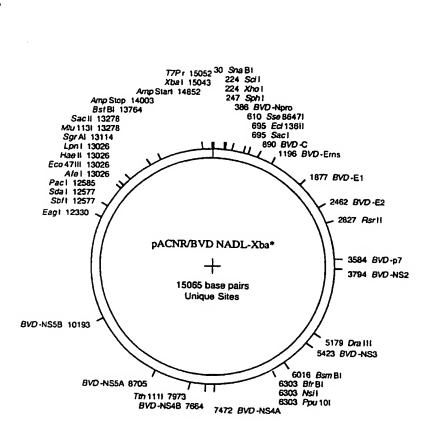


FIGURE 9

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pACNR/BVD NADL-Xba* -> Genes

CNA sequence 15065 b.p. gtatacgagaat ... cgactcactata circular

pacnr/BVD NADL-Xba = HaeII and XhoI digest of pacnr/BVD NADL ligated to HaeII and XhoI digest of pacnr1180/DraIII-/BVD5 8/27 corrected nt 12136 G to C to give HpaI site.

Co

.......

1	gtat	acga	gaat	taga	aaag	gcac	tcgt	atac	gtat	tggg	caat	taaa	aaca	ataa	ttag	gcct	aggg	jaaca	aato	cctc	80
81	tcaç	rcgaa	ggcc	gaaa	agag	gcta	igcca	tgcc	ctta	gtag	gact	agca	taat	gagg	3999	tago	aaca	gtgg	rtgag	ttcg	160
161	ttgg	atgg	ctta	agco	ccga	gtac	aggg	tagt	cgto	agtg	gttc	gacg	cctt	ggaa	taaa	ggto	tcga	gatg	jecac	gtgg	240
241	acga	gggc	atgo	ссаа	agca	cato	ttaa	cctg	ageg	9999	tege	ccag	gtaa	aago	agtt	ttaa	ccga	ctg	CACO	jaata	320
321 1	cago	ctga	tagg	gtgc	tgca	gagg	ccca	ctgt	attg	ctac	taaa	aato	tctg	ctgt	acat	.ggca	C AT	NG CA	C TI	rc	394 3
	ATC I				CTT L	TTA L	TAC Y	AAA K	ACA T					CCC P	GTC V	GGG G		gag E	GAA E	CCT P	454 23
455 24	GTT V	TAT Y	GAT D		GCA A	GCT G	GAT D	CCC P	TTA L		COT G				GCA A	GTC V	CAC H	CCT P	CAA	TCG S	514 43
515 44						CAC H	AAG K							CCA P	ACC T	AAC N	TTG L	GCA A	TCC S	TTA L	574 63
575 64		AAA K			GAC D	TGC C	AGG R	TCG S		AAT N			GGA G	CCT P	GTG V	AGC S	GGG G	ATC I	TAC Y	CTG L	634 83
635 84		CCA P	GGG G	CCA P	CTA L	TTT F	TAC Y	CAG Q	GAC D	TAT Y		GGT G		GTC V	TAT Y	CAC H	AGG R	GCC A	CCG P	CTG L	694 103
695 104		CTC L	TTT F	GAG E	GAG E	GGA G	TCC S	ATG M	TGT C	GAA E	ACG T	ACT T		CCC R	ATA I	GGG G	AGA R	GTA V	ACT T	GGA G	754 123
755 124	AGT S	GAC D	GGA G	AAG K	CTG L	TAC Y	CAC H	ATT I	TAT Y	org C	TGT C	ATA I		GGA G	TGT C	ATA I	ATA I	ATA I	AAA K	AGT S	814 143
815 144		ACG T	AGA R	AGT S	TAC Y	CAA Q	AGG R	GTG V	TTC F	AGG R	TGG W	GTC V	CAT H	AAT N	AGG R	CTT L	GAC D	TGC	CCT P	CTA L	874 163
875 164		GTC V	ACA T	ACT T	TGC	TCA S	GAC D	ACG T	AAA K	GAA E			GCA A	ACA T		AAG K	AAA K	ACA T	CAG Q	AAA K	934 183
935 184		GAC D	AGA R	CTA L	GAA E	AGG R	GGG G	AAA K	ATG M	AAA K	ATA I	GTG V	CCC P	AAA K	GAA E	TCT S	GAA E	AAA K	GAC	AGC S	994 203
995 204		ACT T	AAA K	CCT P	CCG P	GAT D	GCT A	ACA T	ATA I	GTG V	CTC V	GAA E	GGA G	GTC V	AAA K	TAC Y	CAG Q	GTG V	AGG R	AAG K	1054 223
1055 224		GGA G	AAA K	ACC T	AAG K	AGT S	AAA K	AAC N	ACT T	CAG Q	GAC D	GGC	TTG L	TAC Y	CAT H	AAC N	AAA K	AAC N	AAA K	CCT P	1114 243
1115 244		GAA E	TCA S	CGC R	AAG K	AAA K	CTG L	GAA E	AAA K	GCA A	TTG L	TTG L	GCG	TGG W	GCA A	ATA I	ATA I	GCT A	ATA I	GTT V	1174 263
1175 264		TTT F	CAA Q	GTT V	ACA T	ATG M	GGA G	GAA E	AAC N	ATA I	ACA T	CAG Q	TOG	AAC N	CTA L	CAA	GAT D	AAT N	GGG G	ACG T	1234 283
1235 284		GGG G	ATA I	CAA Q	CCGC R	GCA A	ATG M	TTC F	CAA Q	AGG R	OGT G	CTC V	AAT N	AGA R	AGT S	TTA L	CAT H	GGA G	ATC	TCC W	1294 303
1295 304	CCA P	GAG E	AAA K	ATC I	TGT C	ACT T	OCT G	GTC V	CCT P	TCC S	CAT H	CTA L	GCC	ACC T	GAT D	ATA I	GAA E	CTA L	AAA K	ACA T	1354 323
1355 324	ATT	CAT H	GCT G	ATG M	ATG M	GAT D	GCA A	AGT S	GAG E	AAG K	ACC T	AAC N	TAC Y	ACG T	TGT	TGC	AGA R	CTT L	CAA	CGC R	1414 343
1415	CAT	GAG E	TGG W	AAC N	AAG K	CAT H	CCT	TGG W	TGC C	AAC N	TGG W	TAC	AAT N	ATT I	GAA E	CCC	TGG W	ATT	CTA L	GTC	1474 363
1479 364		AAT N	AGA R	ACC	CAA	GCC	AAT N	CTC	ACT	GAG E	GGA G	CAA	CCA P	CCA P	AGG R	GAG E	TGC	GCA A	_	ACT T	1534 383
1535 384	TCT	AGG R	TAT	GAT D	AGG R	GCT A	AGT S	GAC	TTA L	AAC N	CTC	GTA V	ACA T	CAA	GCT A	AGA R	GAT D	AGC S	CCC P	ACA T	1594 403
1599	ccc	TTA	ACA	. ccr	TGC	AAC	AAA	GGA	AAG	AAC	TTC	TCC	TTT	•	GGC	ATA	•	ATG	CGG	•	1654
404		L	T	G	С	K	K	G	K	N	F	S	F		G.	7	1.	M	D	Ġ.	423

FIGURE 10-1

1655 424		TGC C	AAC N	TTT F	gaa E	ATA 1	GCT A	GCA A	AGT S	GAT D	GTA V	TTA L	TTC F	AAA K		CAT H	GAA E	CGC R	TTA I	ACT S	1714 443
1715 444		TTC F	CAG Q	GAT D	ACT T	ACT T	CTT L	TAC Y	CTT L	GTT V	GAC D	GGG G	TTG L	ACC T	AAC N	TCC S	TTA L	GAA E	CCT G	GCC A	1774 463
1775 464		CAA Q	GGA G	ACC T	GCT A	AAA K	CTG L	ACA T	ACC T	TGG W	TTA L	GGC G	aag K	CAG Q	CTC L	GGG	ATA 1	CTA L		AAA K	1834 483
1835 484		TTG L	GAA E	AAC N	AAG K	AGT S	aag K		TGG W	TTT F	GGA G	GCA A	TAC Y	GCT A	GCT A	TCC S	CCT P	TAC Y	TGT C	GAT D	1894 503
1895 504	GTC V	GAT D	CGC R	AAA K	ATT I	GCC G	TAC Y	ATA I	TGG W	TAT Y	ACA T	AAA K	AAT N	TGC C	ACC T	CCT P	GCC	TGC C	TTA L	CCC P	1954 523
1955 524	AAG K	AAC N	ACA T	AAA K	ATT I	GTC V	GGC	CCT P	GGG G	AAA K	TTT F	GAC D	ACC T	AAT N	GCA A	GAG E	GAC D	GGC G	AAG K	ATA I	2014 543
2015 544	TTA L	CAT H	GAG E	ATG M	966 G	GCT G	CAC H	TTG L	TCG S	GAG E	GTA V	CTA L	CTA L	CTT L	TCT S	TTA L	GTG V	GTG V	CTG L	TCC S	2074 563
2075 564	GAC D	TTC F	GCA A	CCG P	GAA E	ACA T	GCT A	ACT S	GTA V	ATG M	TAC Y	CTA L	ATC I	CTA L	CAT H	TTT F	TCC S	ATC I	CCA P	CAA Q	2134 583
2135 584	AGT S	CAC H	GTT V	GAT D	GTA V	ATG M	GAT D	TGT C	GAT D	AAG K	ACC T	CAG Q	TTG L	AAC N	CTC L	ACA T	GTG V	GAG E	CTG L	ACA T	2194 603
2195 604	ACA T	GCT A	GAA E	GTA V	ATA I	CCA P	GGG G	TCG S	orc orc	TGG W	AAT N	CTA L	GGC G	AAA K	TAT Y	GTA V	TGT C	ATA I	AGA R	CCA P	2254 623
2255 624	AAT	TOG	TGG W	CCT P	TAT Y	GAG E	ACA T	ACT T	GTA V	GTG V	TTG L	GCA A	TTT F	GAA E	GAG E	GTG V	AGC S	CAG Q	GTG V	GTG V	2314 643
2315 644	AAG K	TTA L	GTG V	TTG L	AGG R	GCA A	CTC L	AGA R	GAT D	TTA L	ACA T	CGC R	ATT	TGG W	AAC N	GCT A	GCA A	ACA T	ACT T	ACT T	2374 663
2375 664	GCT	TTT F	TTA L	GTA V	TGC	CIT L	GTT V	AAG K	ATA I	GTC V	AGG R	GGC G	CAG Q	ATG M	GTA V	CAG Q	GGC G	ATT I	CTG L	TGG W	2434 683
2435 684	CTA	CTA L	TTG L	ATA I	ACA T	GGG G	GTA V	CAA	GGG G	CAC H	TTG L	GAT D	TGC C	AAA K	CCT P	GAA E	TTC F	TCG S	TAT Y	GCC A	2494 703
2499 704	ATA	GCA A	AAG K	GAC D	GAA E	AGA R	ATT	GGT G	CAA Q	CTG L	GGG G	GCT A	GAA E	GGC G	CTT L	ACC T	ACC T	ACT T	TGG W	aag K	2554 723
2555	GAA	TAC	TCA S	CCT P	GGA	ATG M	AAG K	CIG L	GAA E	GAC D	ACA T	ATG M	GTC V	ATT I	GCT A	TGG W	TGC	GAA E	GAT D	GGG G	2614 743
	AAG K	TTA L	ATG M	TAC	CTC L	CAA	AGA R	TOC	ACG T	AGA R	GAA E	ACC	AGG R	TAT Y	CTC L	GCA A	ATC	TTG L	CAT H	ACA T	2674 763
	ACA	, occ	TTG L	P CCG	ACC T	AGT S	GTG V	GTA V	TTC F	K	AAA K	CTC L	TTT F	GAT	GGG	CGA R	AAG K	CAA Q	GAG E	GAT D	2734 783
	OTA 4 V	ु ए	GAA E	ATG	AAC N	GAC D	AAC N	TTT F	GAA	TTT	GGA	CTC L	TOC	CCA P	TGT C	GAT D	GCC	AAA K	CCC P	ATA I	2794 803
	5 OTA	AGA R	G	AAC K	TTC F	TAA :	ACA T	ACG T	CTG L	CTG L	AAC N	GGA G	CCG	GCC	TTC	CAG Q	ATG M	GTA V	TGC C	CCC P	2854 823
	5 ATA 4 I	G GC	TGC W	ACA T	G G	ACT T	GTA V	AGC S	TGI C	T ACG	TCA S	TTC	AAT N	ATG M	GAC D	ACC	TTA L	GCC A	ACA T	ACT T	2914 843
	5 GT(4 V	GT)	A COC	T AC	TAT Y	' AGA	AGG R	TCT S	AAA K	CCA P	TTC F	CCT P	CAT H	AGG R	CAA	GGC	TGT C	ATC	ACC	CAA	2974 863
	5 AAC 4 K	AAT N	ר כדת נ	G 000	GAC E	GA1	CTC	CAT H	AAC N	TGC C	ATC I	cm L	G GGA	GGA G	AAT N	W TGG	ACT T	C 1C1	CTC V	CCT P	3034 883
	5 GG/ 4 G															TGG W			TAT Y	CAA	3094 903
	5 TF		A GA																AAC N		3154 923
	5 AC 4 T	r oc	T TAC Y	R AGK	CT/	V GT/	GAC D	AGT S	ACC T	TCT S	r rcc c	AAT N	R AGA	GAJ E		v GTG	GCC A	ATA I	GTA V	CCA P	3214 943
		A OG		A TT	A AAC		: AAC K			K			r GT/	CAC Q	GTC V	E ATA	A GC1	ATC M	GAT D	, VCC	3274 963
327 96	5 AA 4 K	A CT	C GG	A CC	T ATY	CC.	, 100 C	R AG	CC/ P	TAT Y	GAJ E	I	I AT	TC/ S	A AGT	CAC E	G G	3 CC7	CTA V	GAA E	3334 983
	5 AA	G AC	A GC			r TN														AGA	3394 1003

3395 GAC AGC TAC TTT CAG CAA TAC ATG CTA AAA GGA GAG TAT CAA TAC TGG TTT GAC CTG GAG 1004 D S Y F Q Q Y M L K G E Y Q Y W F D I. F 3455 GTG ACT GAC CAT CAC CGG GAT TAC TTC CCT GAG TCC ATA TTA GTG GTG GTA GTA GCC CTC 1024 V T D H H R D Y F A E S I L V V V V A L 3515 TTG GGT GGC AGA TAT GTA CTT TGG TTA CTG GTT ACA TAC ATG GTC TTA TCA GAA CAG AAG 1044 L G G R Y V L W L L V T Y N V L S E Q K 3574 1063 3575 GCC TTA GGG ATT CAG TAT GGA TCA GGG GAA GTG GTG ATG ATG GGC AAC TTG CTA ACC CAT 1064 A L G I Q Y G S G E V V M M G N L L T H 3634 3635 AAC AAT ATT GAA GTG GTG ACA TAC TTC TTG CTG CTG TAC CTG CTG AGG GAG GAG AGC 1084 N N I E V V T Y F L L L Y L L R E E S 1103 3695 GTA AAG AAG TGG GTC TTA CTC TTA TAC CAC ATC TTA GTG GTA CAC CCA ATC AAA TCT GTA 3754 3755 ATT GTG ATC CTA CTG ATG ATT GGG GAT GTG GTA AAG GCC GAT TCA GGG GGC CAA GAG TAC 1124 I V I L L M I G G D V V K A D S G G G C CAA GAG TAC 3814 3815 TTG GGG AAA ATA GAC CTC TGT TTT ACA ACA GTA GTA CTA ATC GTC ATA GGT TTA ATC ATA 1144 L G K I D L C F T T V V L I V I G L I I 3874 3875 GCC AGG CGT GAC CCA ACT ATA GTG CCA CTG GTA ACA ATA ATG GCA GCA CTG AGG GTC ACT 1164 A R R D P T I V P L V T I H A A L R V T 3934 1183 3935 GAA CTG ACC CAC CAG CCT GGA GTT GAC ATC GCT GTG GCG GTC ATG ACT ATA ACC CTA CTG 1184 E L T H Q P G V D I A V A V M T I T L L 3994 4054 4055 CTG GTA TCT GCG GTG TTC TTG ATA AGA AGC CTA ATA TAC CTA GGT AGA ATC GAG ATG CCA 1224 L V S A V F L I R S L I Y L G R I E M P 4115 GAG GTA ACT ATC CCA AAC TGG AGA CCA CTA ACT TTA ATA CTA TTA TAT TTG ATC TCA ACA 1244 E V T I P N W R P L T L I L L Y L I S T 4174 4175 ACA ATT GTA ACG AGG TGG AAG GTT GAC GTG GCT GGC CTA TTG TTG CAA TGT GTG CCT ATC 1264 T I V T R W K V D V A G L L L Q C V P I 4234 1283 4235 TTA TTG CTG GTC ACA ACC TTG TGG GCC GAC TTC TTA ACC CTA ATA CTG ATC CTG CCT ACC 1284 L L L V T T L W A D F L T L I L P T 4294 4295 TAT GAA TTG GTT AAA TTA TAC TAT CTG AAA ACT GTT AGG ACT GAT ATA GAA AGA AGT TGG 1304 Y E L V K L Y Y L K T V R T D I E R S W 4354 1323 4355 CTA GOG GGG ATA GAC TAT ACA AGA GTT GAC TCC ATC TAC GAC GTT GAT GAG AGT GGA GAG 1324 L G G I D Y T R V D S I Y D V D E S G E 4414 4415 GGC GTA TAT CTT TTT CCA TCA AGG CAG AAA GCA CAG GGG AAT TTT TCT ATA CTC TTG CCC 1344 G V Y L F P S R Q K A Q G N F S I L L P 4474 1363 4475 CTT ATC AAA GCA ACA CTG ATA AGT TGC GTC AGC AGT AAA TGG CAG CTA ATA TAC ATG AGT 1364 L I K A T L I S C V S S K W Q L I Y M S 4534 4535 TAC TTA ACT TTG GAC TTT ATG TAC TAC ATG CAC AGG AAA GTT ATA GAA GAG ATC TCA GGA 4594 4595 GGT ACC AAC ATA ATA ATA TCC AGG TTA GTG GCA GCA CTC ATA GAG CTG AAC TGG TCC ATG GAA 1404 G T N I I S R L V A A L I E L N W S H E 4655 GAA GAG GAG AGC AAA GGC TTA AAG AAG TTT TAT CTA TTG TCT GGA AGG TTG AGA AAC CTA 4714 4715 ATA ATA AAA CAT AAG GTA AGG AAT GAG ACC GTG GCT TCT TGG TAC GGG GAG GAA GTC 1444 I I K H K V R N E T V A S W Y G E E E V 4774 4775 TAC GGT ATG CCA AAG ATC ATG ACT ATA ATC AAG GCC AGT ACA CTG AGT AAG AGC AGG CAC 1464 Y G H P K I H T I I K A S T L S K S R H 4835 TGC ATA ATA TGC ACT GTA TGT GGG GGC CGA GAG TGG AAA GGT GGC ACC TGC CCA AAA TGT 1484 C I I C T V C E G R E W K G G T C P K C 4894 4895 GGA CGC CAT GGG AAG CCG ATA ACG TCT GGG ATG TCG CTA GCA GAT TTT GAA GAA AGA CAC 1504 G R H G K P I T C G M S L A D F E E R H 4954 5015 AAG CAT AGG AGG TTT GAA ATG GAC CGG GAA CCT AAG AGT GCC AGA TAC TGT GCT GAG TGT 1544 K H R R F E M D R E P K S A R Y C A E C 5074 5075 AAT AGG CTG CAT CCT GCT GAG GAA GGT GAC TTT TGG GCA GAG TCG AGC ATG TTG GGC CTC
1564 N R L H P A E E G D F W A E S S H L G L 5134

FIGURE 10-3

	135 584		ATC I	ACC T	TAC Y	TTT F	GCG A	CTG L	ATG M	GAT D	GGA G	AAG K	ong V	TAT Y	GAT D	ATC I	ACA T	GAG E	TGG W	GCT A	GGA G	5194 1603
	195 604		CAG Q	CGT R	crc v	GGA G	ATC I	TCC S	CCA P	GAT D	ACC T	CAC H	AGA R	GTC V	CCT P	TGT C	CAC H	ATC I	TCA S	TTT F	CCT C	5254 1623
	255 624		CGG R	atg M	CCT P	TTC F	AGG R	CAG Q	GAA E	TAC Y	AAT N	G G G	TTT F	GTA V	CAA Q	TAT Y	ACC T	OCT A	AGG R	GCC	CAA Q	5314 1643
	315 644		TTT F	CTG L	AGA R	AAC N	TTG L	CCC P	GTA V	r CIC	GCA A	ACT T	AAA K	GTA V	AAA K	ATG M	CTC L	atg M	GTA V	e ecc	aac N	5374 1663
	375 664		CGA G	GAA E	GAA E	ATT I	GCT G	AAT N	CTG L	gaa E	CAT H	CTT L	GGG G	TGG W	ATC I	CTA L	AGG R	œc G	CCT P	GCC A	GTG V	5434 1683
_	435 684		AAG K	AAG K	ATC I	ACA T	GAG E	CAC H	GAA E	AAA K	TCC C	CAC H	ATT I	aat N	ATA I	CTG L	GAT D	AAA K	CTA L	ACC T	GCA A	5494 1703
	495 704		TTC F	GGG G	ATC I	ATG M	CCA P	AGG R	GGG G	ACT T	ACA T	b CCC	AGA R	QCC A	CCG P	GTG V	AGG R	TTC F	CCT P	ACG T	AGC S	5554 1723
	555 724		CTA L	AAA K	GTG V	ACG R	AGG R	GCT G	CIG L	gag E	ACT T	GCC A	TGG W	GCT À	TAC Y	ACA T	CAC H	CAA Q	GCC G	996 G	ATA I	5614 1743
	615 744		TCA S	GTC V	GAC D	CAT H	GTA V	ACC T	GCC A	GGA G	AAA K	GAT D	CTA L	CTG L	GTC V	TGT C	GAC D	AGC S	ATG H	GGA G	CGA R	5674 1763
	675 764		AGA R	GTG V	GTT V	TGC C	ÇAA	AGC S	AAC N	AAC N	AGG R	TTG L	ACC T	GAT D	GAG E	ACA T	gag E	TAT Y	GGC G	GTC V	aag K	5734 1783
	735 784		GAC D	TCA S	GGG G	TGC	CCA P	GAC D	oct G	Y CCC	AGA R	TGT C	TAT Y	GTG V	TTA L	AAT N	CCA P	gag E	GCC A	ਹਾ ਪ	AAC N	5794 1803
	795 804		TCA S	GGA G	TCC S	AAA K	GGG G	GCA A	GTC V	GTT V	CAC H	CTC L	CAA Q	AAG K	ACA T	GGT G	GGA G	gaa E	TTC F	ACG T	TGT C	5854 1823
	855 1824		ACC T	GCA A	TCA S	GGC	ACA T	CCG P	GCT A	TTC P	TTC F	GAC D	CTA L	AAA K	AAC N	TTG L	AAA K	GGA G	TGG W	TCA S	GGC G	5914 1843
	5915 1844		P CCT	I I	F	E GYY	GCC A	TCC S	AGC S	GGG	AGG R	GTG V	GTT V	c ccc	AGA R	orc orc	AAA K	GTA V	GGG G	AAG K	AAT N	5974 1863
	5975 1864		GAG E	TCI S	K	P	ACA T	K	ATA I	ATG M	AGT S	GCA G	ATC I	CAG Q	ACC T	GTC V	TCA S	K	AAC N	AGA R	GCA A	6034 1883
	6035 1884		L L	ACC T	GAG E	ATG M	onc v	AAG K	AAG K	ATA I	ACC T	AGC S	ATG M	N N	AGG R	G G	GAC D	TTC F	AAG K	CAG Q	ATT I	6094 1903
	6095 1904		TTC L	GCA A	ACA T	G G	GCA A	G G	K K	ACC	ACA T	GAA E	CTC L	P CCA	K K	GCA A	CTT V	ATA I	GAG E	GAG E	ATA I	6154 1923
	6155 1924		AG/ R	CAC H	: AAC	AGA R	GTA V	TTA L	GTI V	CT1	I I	CCA P	TTA L	AGC R	GCA A	GCG A	GCA A	GAG E	TCA S	GTC V	TAC Y	6214 1943
	6215 1944		TAT Y	PTA 1	AGA R	L TTC	K K	CAC H	P	AGC S	ATC I	TCT S	F	N N	CTA L	AGG R	ATA I	GGG	GAC D	ATG M	AAA K	6274 1963
	6275 1964		G G	GAC D	ATC H	GC/ A	ACC T	G G G	I I	ACC T	TAT Y	A GCA	TC)	TAC Y	G G	Y	TTC F	TGC	CAA	ATG M	CCT P	6334 1983
	633 <u>9</u> 1984		P CC	A AAC	CTY L	R AG	A GCT) A	M M	V V	GA.	Y Y	TC# S	Y	AT#	P TTC	TTA L	GAT D	E GAA	TAC Y	CAT H	6394 2003
	6399 200		P GC	T AC	r cc	E GA	Q CAJ	L CTC	y Y	I	I I	G G G	K K	TA E	CAC H	R AGA	F TTT	TC)	GAG E	AGT S	I	6454 2023
	645! 202		G GT	r Gr	G GC	M S	T ACT	A CCC	T ACC	P P	A GC/	G	S TCC	S GTY	T ACC	T AC	T	G	Q CAV	K	CAC H	6514 2043
		5 CC.				A TT	I	A GCC	P	GA(S GALY	M M	S AAI K		G GAG	G GA1	r CTI		AG1	Q CAG	TTC F	6574 2063
	657 206	5 CT	T GA	T AT	A GC	A GG	C TT.	A AAJ	A AT	P CC	A GTC		F GA		G AA		N N	M M	L TTC	v V	TTT P	6634 2083
	208	4 V	P	T	R	N	M	A	V	Ξ	V	A	K	K	Ĺ	K	A	Κ.	G	Y	N AAC	6694 2103
		5 TC 4 S		A TA Y	C TA	T TA	C AG	T GG.	A GA	G GA	T CC. P	A GC	C AA'	T CT	G AG R	A GT	r GTK	T AC	S TC	, cv	TCC S	6754 2123
		5 CC 4 P		T GI			G GC		A AA N	T GC						G AC T		P CC		r Tro	GAC D	6814 2143
																						6874 2163

PCT/US99/08850

8615 2744	AGG R						TTA L	TTC F	ACA T	TTG L	ATA I	ATG M	ተገተ F	gaa e	GCC	TTC F	GAG E	TTA L	TTA L	ogg G	8674 2763
8675 2764	ATG H	GAC D			GGG G		ATA I	agg R	aac N	r CIG	TCC S	GGA G	AAT N	TAC Y	ATT I	TTG L	GAT D	TTG L	ATA I	TAC Y	8734 2783
8735 2784	GCC		CAC H		CAA Q	ATC I	AAC N	AGA R	GGC G	CTG L	AAG K	AAA K	atc M	GTA V	CTG L	GGC G	TGG W	GCC A	CCT P	GCA A	B794 2803
8795 2804	CCC P	777 F	agt S	TCT C	GAC D	TGG W	ACC T		agt S	GAC D					TTG L	CCA P				TAT Y	8854 2823
8855 2824	TTG L	AGG R	GTA V	GAA E	ACC T	AGG R	TGC C	CCA P	TOT C	GCC G	TAT Y	GAG E	ATG M	AAA K	GCT A	TTC F	AAA K	AAT N	GTA V	G CT G	8914 2843
8915 2844	GGC G	AAA K	CTT L	ACC T	AAA K	GTG V	gag E	gag E	acc S	G G				TGT C			AGA R			agg R	8974 2863
8975 2864	GGA G	CCA P	GTC V	AAC N	TAC Y	AGA R	v esc	ACC T	AAG K	TAT Y	TAC Y	GAT D	GAC D	AAC N	CTC L	aga R	GAG E	ATA I	AAA K	CCA P	9034 2883
9035 2884	GTA V	GCA A	AAG K	TTG L	GAA E	GGA G	CAG Q	CTA V	gag E	CAC H	TAC Y	TAC Y	AAA K	GGG G	GTC V	ACA T	GCA A	AAA K	ATT I	GAC D	9094 2903
9095 2 90 4	TAC Y	act S	AAA K	OGA G	AAA K	ATG H	CTC L	TTG L	GCC A	ACT T	GAC D	AAG K	TGG W	GAG E	GTG V	gaa E	CAT H	GCT C	GTC V	ATA I	9154 2923
2924		R	L	A	K	R	Y	T	G	V	G	F	N	G	Α	Y	L	G	D	E	9214 2943
9215 2944	P	AAT N	CAC H	CGT R	OCT A	CTA L	GTG V	GAG E	AGG R	GAC D	TGT C	GCA A	ACT T	ATA I	ACC T	AAA K	AAC N	ACA T	GTA V	CAG Q	9274 2963
9275 2964	TTT F	CTA L	K	ATG M	AAG K	AAG K	GCIG G	TGT C	GCG A	TTC F	ACC T	TAT Y	GAC D	CTG L	ACC T	ATC I	TCC S	aat N	r CIG	ACC T	9334 2983
2984		L	1	E	L	V	н	R	N	N	L	E	E	K	E	I	P	T	A	T	9394 3003
3004		T	T	W	L	A	Y	T	F	V	N	E	D	V	G	T	I	K	P	V	9454 3023
9455 302	S CTA	G G	GAG E	AGA R	GTA V	ATC I	P	GAC D	P	gta V	CTT V	GAT D	ATC I	AAT N	TTA L		P CCY	GAG E	CTG V	CAA Q	9514 3043
951: 304	5 GTG 4 V	GAC D	T ACG	TCA S	GAG E	CTT V	C	ATC I	ACA T	ATA I	ATT I	G G	AGG R	gaa E	ACC T	CTG L	ATG M	ACA T	ACG T	GGA G	9574 3063
957 306	s ond	T ACA	P	v crc	TTG L	GAA E	K	GTA V	GAG E	CCT P	GAC D	GCC A	AGC S	GAC D	N N	CVY	AAC N	TCG S	GTG V	AAG K	9634 3083
963 308	5 ATC	G G	L	GAT D	GAG E	G	N N	TAC	P	G	P	GGA G	ATA I	CAG Q	ACA T	CAT H	ACA T	CTA L	ACA T	GAA E	9694 3103
310		I	Н	N	R	D	A	R	P	F	I	М	I	L	G	S	R	N	S	I	9754 3123
312		N	R	A	K	т	A	R	N	I	N	L	Y	T	G	N	D	P	R	E	9814 3143
314	5 AT/ 4 I	R	D	L	M	λ	A	G	R	M	L	V	V	A	L	R	D	٧	D	P	9874 3163
	5 GX 4 E	CTC L	S TCT	GA/ E	M M	y V	GAT D	F	K AAC	G	T ACT	F	L	GAT D	AGG R	E GAG	A A	CTG L	gag E	GCT A	9934 3183
318	5 CT/	A AG	ר כזא	: GGC	CN	v cci	CAA 1	coc	, AAC	CAC	: CTT	ACC	AAG	GAN	GCT	CIT	AGG	AAT	TTG	ATA	9994 3203
000	4 L	s			Q	P		P	K	Q	V	т	ĸ	E	A	٧	R	N	L	•	
320	4 L 5 GA 4 E	A CAC	G AA	K	Q A GAT D	v V	GAC E	I	P	N	W	F	A	s	D	D	P	V	F	L	10054 3223
1005 322	5 GA 4 E 5 GA 14 E	A CAC	G GCC	K TT/ L	Q A GAT D A AA K	CTC V A AA1	G GAC E C GA1 D	I K	P TAC Y	N TAC Y	W TTA	F GTA V	A GGA G	S GAT D	O CTT	D GGA G	GAG E	CTA L	F AAA K	L GAT D	10054 3223 10114 3243
1005 322 1011 324	5 GA 4 E 5 GA 4 E 5 GA 14 E	A CAI Q A GTV A GC A	G GCC A	A GC:	Q GAT D A AAA K	CON	G GAC E T GAT D G GCC A	I AAC K ACC T	P TAC Y G GA' D	N TAC Y CAC O	W L L S ACA	F GTA V AGA R	A G G ATT	S GAT D ATA	D GTT	D G G GAC E	GAG E GTA	V CTA GGG	F AAA K TCA S	GAT D AGG R	10054 3223 10114 3243 10174 3263
1005 322 1011 324 1017 326	14 L 15 GAU 14 E 15 GA 14 E 15 CA 14 Q 175 AC	A CAC Q A GTN A GC A GC Y	G GCC A T AA K T GCC A	A GC	Q GAT R CT L K	CTC G	G GAC E T GAN D G GCC A	I AAC K	P TAC Y G GA' D T TGC	N TAC Y CAC Q TTC F	W L E AC/ T CTX L	F GTA V AGA R	A GGA G ATTI	S GAT D T AT/ I T TC/ S	D GTT V AAG K AAG	D G G G G G G G K	GAG E GTA V CAG	CTA L GGG G ATC	F AAA K TCA S AGT S	L GAT D AGG R	10054 3223 10114 3243 10174 3263 10234 3283
320 1005 322 1011 324 1017 326 1023 328	14 L 15 GA 14 E 15 GA 14 C 15 AC 15 AC 15 AC 15 AC	A CAC Q A GTA A G TA T CC	G AAAA K T GC A A CT A	A GCATT	Q GAN K	CT. GGC	G GCC A A TCT S	I AAC K T AGC S TTV	P TAC Y GAT D TGC W GCT L	N COX	W E TTA L G ACA T CTC L CTC C	F GTA V AGA R AAC K	A GGA	S GAT D T AT? T TC! S	D GTT V AAG	D GGAGE RAAGE K	GAGGE CAGG	CTA L GGC G ATC H	F AAA K S AGT S AAC K	L GAT D AGG R	10054 3223 10114 3243 10174 3263 10234

1	0355 3324		CTA L	GGT G	ACA T	ATA I	CCA P	GCC A	AGA R	AGG R	V GIG	AAG K	ATA I	CAC H	CCA P	TAT Y	GAA E	GCT A	TAC Y	CTC	AAG K	10414 3343
3	0415 3344		AAA K	GAT D	TTC F	ATA I	gaa E	GAA E	gaa E	GAG E	AAG K	AAA K	CCT P	AGG R	GTT V	AAG K	GAT D	ACA T	GTA V	ATA	AGA R	10474 3363
1	0475 3364		CAC H	AAC N	AAA K	TGG W	ATA I	CTT L	AAA K	AAA K	ATA I	AGG R	TTT F	CAA Q	GGA G	AAC N	CTC L	AAC N	ACC T	AAG K	AAA K	10534 3383
1	0535 3384		CTC L	AAC N	CCG P	GGG G	AAA K	CTA L	TCT S	gaa E	CAG Q	TTG L	GAC D	AGG R	GAG E	OGG G	CGC R	AAG K	AGG R	AAC N	ATC I	10594 3403
1	0595 3404	TAC Y	AAC N	CAC H	CAG Q	TTA I	GCT G	ACT T	ATA I	ATG M	TCA S	AGT S	GCA A	GGC G	ATA I	AGG R	CTG L	GAG E	AAA K	TIG L	CCA P	10654 3423
1	0655 3424	ATA I	GTG V	AGG R	GCC A	CAA Q	ACC T	GAC D	ACC T	AAA K	ACC T	777 F	CAT H	GAG E	GCA A	ATA I	AGA R	GAT D	AAG K	ATA I	GAC D	10714 3443
1	0715 3444		AGT S	GAA E	AAC N	CGG R	CAA O	aat N	CCA P	gaa E	TTG L	CAC H	AAC N	AAA K	TTG L	TTG L	GAG E	ATT I	TTC P	CAC H	ACG T	10774 3463
1	10775 3464		GCC A	CAA Q	CCC P	ACC T	CTG L	AAA K	CAC H	ACC T	TAC Y	GCT G	GAG E	GTG V	ACG T	TGG W	GAG E	CAA Q	CTT L	GAG E	GCG A	10834 3483
3	0835 3484	GGG G	ATA I	AAT N	AGA R	AAG K	GGG G	GCA A	GCA A	GCC	TTC F	CTG L	GAG E	AAG K	AAG K	AAC N	ATC I	GGA G	GAA E	GTA V	TTG L	10894 3503
1	0895 3504		TCA S	GAA E	AAG K	CAC H	CTG L	GTA V	GAA E	CAA Q	TTG L	GTC V	AGG R	GAT D	CTG L	AAG K	GCC	936 G	AGA R	AAG K	ATA I	10954 3523
1	10955 3 524	AAA K	TAT Y	TAT Y	GAA E	ACT T	GCA A	ATA I	CCA P	AAA K	AAT N	gag e	AAG K	AGA R	GAT D	GTC V	AGT S	GAT D	GAC D	TGG W	CAG Q	11014 3543
3	11015 3544	GCA A	GGC G	GAC D	CTG L	CTC V	GTT V	GAG E	AAG K	AGG R	CCA P	AGA R	GTT V	ATC I	CAA Q	TAC Y	CCT P	GAA E	GCC	AAG K	ACA T	11074 3563
1	11075 3564		CTA L	GCC A	ATC	ACT T	aag K	GTC V	ATG M	TAT Y	AAC N	TGG W	CTC CTC	XXX K	CAG Q	CAG Q	CCC P	GTT V	org O	ATT I	CCA P	11134 3583
:	11135 3584	GGA G	TAT Y	GAA E	GGA G	AAG K	ACC T	CCC P	TTG L	TTC P	AAC N	ATC I	TTT F	GAT D	AAA K	GTG V	aga R	AAG K	GAA E	TGG W	GAC D	11194 3603
	11195 3604	TCG S	TTC F	AAT N	GAG E	CCA P	GTG V	GCC A	GTA V	agt S	TTT F	GAC D	ACC T	AAA K	QCC GCC	TGG W	GAC D	ACT T	CAA Q	A CLC	ACT T	11254 3623
:	11255 3624		AAG K	GAT D	CIG CIG	CAA Q	CTT L	att I	GGA G	GAA E	ATC I	CAG Q	XXX K	TAT Y	TAC Y	TAT Y	AAG K	AAG K	GAG E	TGG W	CAC H	11314 3643
	11315 3644		TTC F	ATT I	GAC D	ACC T	ATC I	ACC T	GAC D	CAC H	ATG M	ACA T	gaa E	GTA V	CCA P	GTT V	ATA I	ACA T	GCA A	GAT D	GCT G	11374 3663
	11375 3664	GAA E	GTA V	TAT Y	ATA I	AGA R	aat N	GGG G	CAG Q	aga R	G G	AGC S	GOC G	CAG Q	CCA P	GAC D	ACA T	act S	GCT A	GGC G	AAC N	11434 3683
	11435 3684	AGC S	ATG M	TTA L	AAT N	GTC V	CTG L	ACA T	ATG M	atg M	TAC Y	GGC	TTC F	TGC C	GAA E	AGC S	ACA T	œc G	GTA V	CCG P	TAC Y	11494 3703
	11495 3704	AAG K	AGT S	TTC F	AAC N	AGG R	v GTG	GCA A	AGG R	ATC I	CAC H	GTC V	TOT C	CCC C	GAT D	CAT D	G G	TTC F	TTA L	ATA I	ACT T	11554 3723
	11555 3724	gaa E	AAA K	GOG G	TTA L	GGC	CTG L	AAA K	TTT F	y GCI	AAC N	AAA K	GGG G	atg M	CAG Q	ATT	CTT L	CAT H	GAA E	y CCY	GCC G	11614 3743
	11615 3744	K	P	Q	ĸ	1	T	E	G	E	ĸ	М	K	V	A	Y	R	F	E	D	1	11674 3763
	11675 3764	GAG E	TTC F	TGT C	TCT S	CAT H	ACC T	CCA P	GTC V	P	v GTT	AGG R	TGG W	TCC S	GAC D	AAC N	ACC T	agt S	AGT S	CAC H	ATG H	11734 3783
	11735 3784	A	G	R	D	T	A	V	I	L	S	K	М	Α	T	R	L	D	S	s	G	11794 3803
	11795 3804	GAG E	AGG R	G	ACC T	ACA T	GCA A	TAT Y	GAA E	AAA K	GCC A	GTA V	y ccc	TTC F	agt S	TTC F	TTG L	CTG L	ATG M	TAT Y	TCC S	11854 3823
	11855 3824	₩	N	P	L	٧	ĸ	ĸ	1	C	L	L	v	L	5	Q	Q	P	E	T	D	11914 3843
	11915 3844	P	S	K	н	^	Т	Y	Y	Y	K	G	D	P	I	G	λ	Y	K	D	V	11974 3863
	11975 3864	1	G	R	N	L	S	E	L	K	R	Т	G	F	Ε	K	L	A	N	L	N	12034 3883
	12035	CIA	AGC	CTG	TCC	ACG	TTG	GGG	ATC	TOG	ACT	AAG	CAC	ACA	AGC	AAA	AGA	ATA	ATT	CAG	GAC	12094

FIGURE 10-7

2095 3904		crr v	GCC A	ATT I	GGG G	AAA K			GGC G	AAC N	TCG W	CTA L	GTT V	AAC N	OCC A	GAC D	AGG R	CTG L	ATA I	TCC S	12154 3923	
12155 3924		AAA K	ACT T	GCC G	CAC H	TTA L	TAC Y	ATA I	CCT P	GAT D	AAA K	GGC G	TTT F	ACA T	TTA L	CAA Q	GGA G	AAG K	CAT H	TAT Y	12214 3943	
12215 3944		CAA Q	CTG L	CAG Q	CTA L	AGA R	ACA T	GAG E	ACA T	AAC N	CCG P	GTC V	ATG M	GGG G	GTT V	GGG G	ACT T	gag E	AGA R	TAC Y	12274 3963	
12275 3964		TTA L	GGT G	CCC P	ATA I	GTC V	AAT N	CTG L	CTG L	CTG L	aga R	AGG R	TTG L	AAA K	ATT I	r cig	CTC L	ATG M	ACG T	GCC A	12334 3983	
12335 3984			GTC V	AGC S	AGC S	TGA	gaca	aaat	gtat	ata	ttgt	aat	aati	taato	cate	jtac a	atagi	tgtal	tata	acat	12408 3989	
12409	agti	cggga	accg	tcca	cctca	aaga	gac	jaca	gcc	aaca	acgc	acag	ctaa	acag	tagto	aaga	tta	tcta	cctc	aagat	12488	
12489	aac	acta	catt	taat	gcaca	acago	cacti	tage	ctgt	atga	ggat	cgc	cega	cgtc	tata	jttg	gac t	ag g g	aaga	cctct	12568	
12569	aac	agcc	CCCL	gcag	gtta	atta	acta	gtgg	gaat	acgc:	9999	catg	cege	gttt	tage	atati	tgac	gacc	CAAL	tctca	12648	
12649	cgt	ttga	cage	ctat	cato	gtcg	agca	agac	gttt	cccg	ttga	atat	ggct	cata	acac	ccct	tgta	ttac	tgtt	tatgt	12728	
12729	aag	caga	cagt	ttta	ttgt	tcat	gatg	atat	attt	ttat	cttg	tgca	atgt.	aaca	tcag	agat	tttg	agac	acgt	ggctt	12808	
12809	tgt	tgaa	taaa	tega	actt	ttgc	tgag	ttga	agga	tcag	atca	cgca	tctt	cccg	acaa	cgca	gaco	gttc	cgtg	gcaaa	12888	
12889	gca	aaag	ttca	aaat	cacc	aact	ggtc	cacc	taca	acaa	agct	ctca	tcaa	ccgt	gget	ccct	cact	ttct	ggct	ggatg	12968	
12969	atg	gggc	gatt	cagg	cctg	gtat	gagt	cagc	aaca	cctt	cttc	acga	ggca	gacc	tcag	cgct	agcg	gagt	gtat	actgg	13048	
13049	ctt	acta	tgtt	ggca	ctga	tgag	ggcg	tcag	tgaa	gtgc	ttca	tgtg	gcag	gaga	aasa	aggc	tgca	ccgg	tgcg	tcagc	13128	
13129	aga	atat	gtga	taca	ggat	atat	tccg	CEEC	ctcg	ctca	ctga	ctcg	ctac	gctc	ggtc	gttc	gact	gcgg	cgag	cggaa	13208	
13209	atg	gctt	acge	recâă	ggcg	gaga	tttc	ctgg	aaga	tgcc	agga	agat	actt	aaca	ggga	agtg	agag	ggcc	acaa	caaag	13288	
13289	ccg	ttt	tcca	tagg	ctcc	gccc	ecct	gaca	agca	tcac	gaaa	tctg	acgo	tcaa	atca	gtgg	tggc	gaaa	cccg	acagg	13368	
13369	act	atas	agat	acca	aggeg	tttc	ccct	ggeg	gcto	ccto	gtgc	gcto	tcct	gttc	ctgc	cttt	cggt	ttac	cggt	gtcat	13448	
13449	tco	gctg	ttat	ggc	gegt	ttgt	ctca	ttee	acgo	ctga	cact	cagt	CCCG	gg ca	ggca	gttc	gctc	caag	ctgg	actgt	13528	
13529	atç	cacq	aaco	cccc	gttc	agto	cgac	cgct	gcgc	ctta	tccg	gtaa	ctat	cgto	ttga	gtec	aacc	cgga	aaga	catge	13608	
13609	aaa	agce	эссво	etgge	agca	igeca	ctgg	taat	tgat	ttag	agga	gtta	gtct	tgaa	gtca	tgcg	ccgg	ttaa	gget	aaact	13688	
13689	gaz	angga	caa	gttt	ggtg	gacto	icg ct	ccto	caag	rccaç	ttac	ctcç	gtto	aaag	agtt	ggta	gctc	agag	aacc	ttcga	13768	
13769	aaa	accç	gece	cgca	aggcg	gtt	ttt	gttt	tcag	agca	agag	jatta	ıcgeç	caga	ccae	aacg	atct	caaç	aaga	tcatc	13848	
13849) tt	scca	aggg	gtcti	gacgo	ccaç	rgga	acga	aaaa	tcac	gtt	aggg	atti	tggt	cato	agat	tato	aaaa	agga	tette	13928	
13929) ac	ctaga	stcc	LLLE	2000	Laaaa	atga	agti	tta	atc	atcı	aaaq	jtati	atate	gagte	MACT	tggt	ctga	cagt	CACCA	14008	
14009	ate	gett	atc.	agtg	aggca	accta	tete	ageg	jatet	gtc	tatti	cgt	cate	CAL	gtt	jectç	jacto	cccç	tcgt	gtaga	14088	
14089	9 ta	BCTA	cgat	acgg	gaggg	gctta	accat	ctg	30000	agt	getge	canto	gata	ccgc	gagad	ccac	gcto	acce	gcto	cagat	14168	
1416	9 tt	atca	gcaa	taaa	ccago	ccago	ccggi	agg	jecga	ageg	caga	gtg	gtcc	tgca	actt	atc	gcct	cca	ccag	tctat	14248	
1424	9 ta	attg	ttgc	cggg	aagc	taga	gtaag	gtagi	ttcg	cag	ttaa	tagt	ttgc	gcaa	egtt	jttg	catt	get	gcag	gcateg	14328	
1432	9 tg:	gtgt	cacg	ctcg	tegt	ttggi	tatgg	ctt	catte	age	teeg	gttc	ccaa	cgat	caag	gcgaq	gttac	catg	atcc	ccatg	14408	
1440	9 tt	gtgc	8888	aagc	ggtt	aget	cctt	ggt	ctc	cgat	cgtt	gtca	gaag	taag	ttgg	ccgc	agtgi	ttat	cacte	catggt	14488	
																					14568	
																					14648	
																					14728	
																					14808	
	-																				14888	
																					14968	
																					15048	
				cact	_	J	9			-								J	-: -	9	15065	
1304	,																					

WO 99/55366 PCT/US99/08850

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BVDV NADL (inf. clone) -> Genes

DNA sequence 12578 b.p. gtatacgagaat ... ctaacagcccc linear

1 gtatacgagaattagaaaaggcactcgtatacgtattgggcaattaaaaataattaggcctagggaacaaatccctc 80 81 tcagcgaaggccgaaaagaggctagccatgcccttagtaggactagcataatgagggggtagcaacagtggtgagttcg 160 161 ttggatggcttaagccctgagtacagggtagtcgtcagtggttcgacgccttggaataaaggtctcgagatgccacgtgg 240 395 ATC ACA AAT GAA CTT TTA TAC AAA ACA TAC AAA CAA AAA CCC GTC GGG GTG GAG GAA CCT 4 I T N E L L Y K T Y K Q K P V G V E E P 455 GTT TAT GAT CAG GCA GGT GAT CCC TTA TTT GGT GAA AGG GGA GCA GTC CAC CCT CAA TCG 515 ACG CTA AAG CTC CCA CAC AAG AGA GGG GAA CGC GAT GTT CCA ACC AAC TTG GCA TCC TTA
44 T L K L P H K R G E R D V P T N L A S L 575 CCA AAA AGA GGT GAC TGC AGG TCG GGT AAT AGC AGA GGA CCT GTG AGC GGG ATC TAC CTG 64 P K R G D C R S G N S R G P V S G 1 Y L 635 AAG CCA GGG CCA CTA TTT TAC CAG GAC TAT AAA GGT CCC GTC TAT CAC AGG GCC CCG CTG 84 K P G P L P Y Q D Y K G P V Y H R A P L 695 GAG CTC TIT GAG GAG GGA TCC ATG TGT GAA ACG ACT AAA COG ATA GGG AGA GTA ACT GGA 104 E L F E E G S M C E T T K R I G R V T G 815 GCC ACG AGA AGT TAC CAA AGG GTG TTC AGG TGG GTC CAT AAT AGG CTT GAC TGC CCT CTA 144 A T R S Y Q R V F R W V H N R L D C P L 874 183 935 CCC GAC AGA CTA GAA AGG GGG AAA ATG AAA ATA GTG CCC AAA GAA TCT GAA AAA GAC AGC 184 P D R L E R G K M K I V P K E S E K D S 995 AAA ACT AAA CCT CCG GAT GCT ACA ATA GTG GTG GAA GGA GTC AAA TAC CAG GTG AAG 204 K T K P P D A T I V V E G V K Y O V R K 1055 AAG GGA AAA ACC AAG AGT AAA AAC ACT CAG GAC GGC TTG TAC CAT AAC AAA AAC AAA CCT 224 K G K T K S K N T Q D G L Y H N K N K P 1114 1115 CAG GAA TCA CGC AAG AAA CTG GAA AAA GCA TTG TTG GCG TGG GCA ATA ATA GCT ATA GTT 244 Q E S. R K K L E K A L L A W A I I A I V 1175 TTG TTT CAA GTT ACA ATG GGA GAA AAC ATA ACA CAG TGG AAC CTA CAA GAT AAT GGG ACG 264 L F Q V T M G E N I T Q W N L Q D N G T 1234 1235 GAA GOG ATA CAA COG GCA ATG TTC CAA AOG GGT GTG AAT AGA AGT TTA CAT GGA ATC TOG 284 E G I Q R A M P Q R G V N R S L H G I W 1295 CCA GAG AAA ATC TGT ACT GGT GTC CCT TCC CAT CTA GCC ACC GAT ATA GAA CTA AAA ACA 1354 1355 ATT CAT GOT ATG ATG GAT GCA AGT GAG AAG ACC AAC TAC ACG TGT TGC AGA CTT CAA CGC 324 I H G H D A S E K T N Y T C C R L Q R 1415 CAT GAG TGG AAC AAG CAT GGT TGG TGC AAC TGG TAC AAT ATT GAA CCC TGG ATT 344 H E W N K H G W C N W Y N I E P W I 1474 363 1475 ATG AAT AGA ACC CAA GCC AAT CTC ACT GAG GGA CAA CCA CCA AGG GAG TCC GCA GTC ACT 364 M N R T Q A N L T E G Q P P R E C A V T 1535 TGT AGG TAT GAT AGG GCT AGT GAC TTA AAC GTG GTA ACA CAA GCT AGA GAT AGC CCC ACA 384 C R Y D R A S D L N V V T Q A R D S P T 1594 403 1595 CCC TTA ACA GGT TGC AAG AAA GGA AAG AAC TTC TCC TTT GCA GGC ATA TTG ATG CGG GGC 404 P L T G C K K G K N F S F A G I L M R G 1655 CCC TGC AAC TTT GAA ATA GCT GCA AGT GAT GTA TTA TTC AAA GAA CAT GAA CGC ATT AGT 1714 1715 ATG TTC CAG GAT ACT ACT CTT TAC CTT GTT GAC GGG TTG ACC AAC TCC TTA GAA GGT GCC 1774

FIGURE 11-2

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4/21/99
BVDV NADL (inf. clone) -> Ge.....
   1775 AGA CAA GGA ACC CCT AAA CTG ACA ACC TGG TTA GGC AAG CAG CTC GGG ATA CTA CGA AAA 464 R O G T A K L T T W L G K O L G I L G K
   1835 AAG TTG GAA AAC AAG AGT AAG ACG TGG TTT GGA GCA TAC GCT GCT TCC CCT TAC TGT GAT 484 K L E N K S K T W F G A Y A A S P Y C D
    1895 GTC GAT CGC AAA ATT GGC TAC ATA TGG TAT ACA AAA AAT TGC ACC CCT GCC TGC TTA CCC
    1955 AAG AAC ACA AAA ATT GTC GGC CCT GGG AAA TTT GAC ACC AAT GCA GAG GAC GGC AAG ATA
                                                                                                                     2014
    2015 TTA CAT GAG ATG GGG GGT CAC TTG TCG GAG GTA CTA CTA CTT TCT TTA GTG GTG CTG TCC 544 L H E M G G H L S E V L L L S L V V L S
                                                                                                                     2074
    2075 GAC TTC GCA CCG GAA ACA GCT AGT GTA ATG TAC CTA ATC CTA CAT TTT TCC ATC CCA CAA
                                                                                                                      583
    2135 AGT CAC GTT GAT GTA ATG GAT TGT GAT AAG ACC CAG TTG AAC CTC ACA GTG GAG CTG ACA
584 S H V D V M D C D K T Q L N L T V E L T
                                                                                                                      2194
    2195 ACA GCT GAA GTA ATA CCA GGG TCG GTC TGG AAT CTA GGC AAA TAT GTA TGT ATA AGA CCA
                                                                                                                     2314
    2255 AAT TOG TOG CCT TAT GAG ACA ACT GTA GTG TTG GCA TTT GAA GAG GTG AGC CAG GTG GTG 674 N W W P Y E T T V V L A F E E V S Q V V
    2315 AAG TTA OTG TTG AGG GCA CTC AGA GAT TTA ACA CGC ATT TGG AAC GCT GCA ACA ACT ACT 644 K L V L R A L R D L T R I W N A A T T T
                                                                                                                      2374
                                                                                                                      663
     2375 GCT TTT TTA GTA TGC CTT GTT AAG ATA GTC AGG GGC CAG ATG GTA CAG GGC ATT CTG TGG 664 A F L V C L V K I V R G Q M V Q G I L W
                                                                                                                      2434
     2435 CTA CTA TTG ATA ACA GOG GTA CAA GOG CAC TTG GAT TGC AAA CCT GAA TTC TCG TAT GCC
                                                                                                                      2494
                                                                                                                      703
     2555 GAA TAC TCA CCT GGA ATG AAG CTG GAA GAC ACA ATG GTC ATT GCT TGC GAA GAT GGG
                                                                                                                      2614
     2615 AAG TTA ATG TAC CTC CAA AGA TGC ACG AGA GAA ACC AGG TAT CTC GCA ATC TTG CAT ACA 744 K L M Y L Q R C T R E T R Y L A I L H T
     2675 AGA GCC TTG CCG ACC AGT GTG GTA TTC AAA AAA CTC TTT GAT GGG CGA AAG CAA GAG GAT 764 R A L P T S V V F K K L F D G R K Q E D
                                                                                                                      2734
     2735 GTA GTC GAA ATG AAC GAC AAC TTT GAA TTT GGA CTC TGC CCA TGT GAT GCC AAA CCC ATA
784 V V E M N D N F E F G L C P C D A K P I
     2795 GTA AGA GGG AAG TTC AAT ACA ACG CTG CTG AAC GGA CCG GCC TTC CAG ATG GTA TGC CCC 804 V R G K F N T T L L N G P A F Q M V C P
                                                                                                                      2854
     2855 ATA GGA TGG ACA GGG ACT GTA AGC TGT ACG TCA TTC AAT ATG GAC ACC TTA GCC ACA ACT R24 I G W T G T V S C T S F N M D T L A T T
     2915 GTG GTA COG ACA TAT AGA AGG TCT AAA CCA TTC CCT CAT AGG CAA GGC TGT ATC ACC CAA 844 V V R T Y R R S K P F P H R O G C I T O
     2975 AAG AAT CTG GGG GAG GAT CTC CAT AAC TGC ATC CTT GGA GGA AAT TGG ACT TGT GTG CCT 864 K N L G E D L H N C I L G G N W T C V P
                                                                                                                      3034
     3035 GGA GAC CAA CTA CTA TAC AAA GGG GGC TCT ATT GAA TCT TGC AAG TGG TGT GGC TAT CAA 884 G D Q L L Y K G G S I E S C K W C G Y Q
                                                                                                                       3094
                                                                                                                       903
     3095 TIT AMA GAG AGT GAG GGA CTA CCA CAC TAC CCC ATT GGC AAG TGT AMA TTG GAG AMC GAG 904 F K E S E G L P H Y P I G K C K L E N E
                                                                                                                      3154
      3155 ACT GOT TAC AGG CTA GTA GAC AGT ACC TCT TOC AAT AGA GAA GGT GTG GCC ATA GTA CCA 924 T C Y R L V D S T S C N R E G V A I V P
                                                                                                                       3214
      3215 CAA GOG ACA TTA AAG TGC AAG ATA GGA AAA ACA ACT GTA CAG GTC ATA GCT ATG GAT ACC 944 O G T L K C K I G K T T V O V I A H D T
      3275 AAA CTC GGA CCT ATG CCT TGC AGA CCA TAT GAA ATC ATA TCA AGT GAG GGG CCT GTA GAA 964 K L G P M P C R P Y E I I S S E G P V E
                                                                                                                       3334
      3335 AAG ACA GCG TGT ACT TTC AAC TAC ACT AAG ACA TTA AAA AAT AAG TAT TTT GAG CCC AGA 984 K T A C T F N Y T K T L K N K Y F E P R
                                                                                                                       3394
      3395 GAC AGC TAC TTT CAG CAA TAC ATG CTA AAA GGA GAG TAT CAA TAC TGG TTT GAC CTG GAG
                                                                                                                        1023
      3455 GTG ACT GAC CAT CAC CGG GAT TAC TTC GCT GAG TCC ATA TTA GTG GTG GTA GTA GCC CTC 1024 V T D H H R D Y F A E S I L V V V V A L
                                                                                                                       3514
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BVDV	NAI	DL	(in	ıf. c	lone) ->	Ge	ر			24	/67									4/2	1/99
3515 1044			T C									CTT V		TAC Y		GTC V			_	_	AAG K	3574 1063
	GCC		A G													GGC G			CTA L	ACC T	CAT H	3634 1083
	AAC N							ACA :		TTC F					CTA L	CTG L				GAG E	AGC S	3694 1103
3699 1104		AA K	G A												GTA V	CAC H			AAA K	TCT S	GTA V	3754 1123
3755 1126		េញ	G A												GAT D	TCA S		GCC G	CAA Q	GAG E	TAC Y	3814 1143
381! 114		G GG														GTC V			TTA L	ATC I	ATA I	3874 1163
		: AG								CCA P						GCA A			AGG R	GTC V	ACT T	3934 1183
393: 118		CT L	7G 7											GCCG A	GTC V	ATG M		ATA I	ACC T	CTA L	CTG L	3994 1203
399 120		3 GT V								TTT F						TTA L		TGC C	att I	CTC L	AGC S	4054 1223
405 122		G GT V														GGT G		ATC I	gag E	ATG H	CCA P	4114 1243
411 124		ទ ទា V											TTA L	ATA I	CTA L	TTA L		TTG L	ATC I	TCA S	ACA T	4174 1263
417 126		I A													TTG L	TTG L		TGT C	GTG V	CCT P	ATC I	4234 1283
423 128		A TI				ACA T					GAC D		TTA L	ACC T	CTA L	ATA I	CTG L	ATC I	CTG L	CCT P	ACC T	4294 1303
	5 TA 4 Y	T GJ				AAA K								AGG R	ACT T	GAT D	ATA I	gaa B	aga R	agt S	TGG W	4354 1323
	5 CT 4 L	A CX G										TCC S				GTT V	GAT D	gag e	AGT S	GGA G	gag E	4414 1343
	5 GG 4 G	c Gi				TTT P	CCA P		AGG R		AAA K		CAG Q	GCG	AAT N	TTT F	TCT S	ATA I	CTC L	TTG L	ECC P	4474 1363
	5 CT 4 L	T AT				ACA T		ATA I			GTC V				W TGG	CAG Q	CTA L	ATA I	TAC Y	ATG M	agt S	4534 1383
	5 TA					GAC D		atg M			ATG M				GTT V	ATA I	gaa E	CAG E	ATC I	TCA S	GCA G	4594 1403
	5 00 4 G	TA		AAC N	ATA I	ATA I	TCC S	AOG R	TTA L	GTG V	GCA A	y CCY	CTC L	ATA I	GAG E	CTG L	AAC N	TGG W	TCC S	ATG M	gaa E	4654 1423
		A G			AGC S	K K	G G		AAG K	AAG K	TTT P	TAT Y	CTA L	TTG	TCT S	GCA G	AGG R	TTG L	AGA R	AAC N	CTA L	4714 1443
144	4 I	I		ĸ	Н	ĸ	v	R	N	E	T	٧	A	S	W	TAC Y	G	E	E	E	V	4774 1463
146	4 Y	G		H	P	ĸ	1	М	T	I	I	K	٨	s	T	L	s	K	s	R	н	4834 1483
140	34 C	I		I	С	T	v	С	E	G	R	Ē	W	K	G	G	T	С	P	K		1503
150	04 G	R	t	н	G	К	P	I	Т	С	G	M	S	L	A	D	P	Е	E	R	H	4954 1523
15	24 Y	K		R	I	F	I	R	Ε	C	N	P	E	G	H	С	s	R	С	Q		1543
15	44 K	Н	ŧ	R	R	F	E	М	D	R	Ε	P	K	s	A	R	Y	С	A	E	C	5074 1563
15	54 N	A	ł	L	Н	P	A	Е	E	G	D	F	W	λ	E	s	s	H	L	G	L	5134 1583
15	84 K	1	1	T	Y	F	A	L	М	D	G	ĸ	V	¥	D	1	т	E	W	A		5194 1603
51 16	95 T	GC (CAG O	CGT R	V CTC	G	I	S	P	GAT D	T	H H	R R	v V	P	r TGT C	H H	I	S TC	F	G	5254 1623

BVD	v 10	JAD	1 6	of.	clone	•) •>	. G.				25/6	7									4/	21/99
52		TCA		ATG	cct	TTC	AGG	CAG	GAA E	TAC	AAT N	GGC	TTT F	GTA V	CAA	TAT Y	ACC		AGG R	GGG G		5314 1643
53		CTA	TTT		AGA	AAC	TTG	ccc	GTA	CTG	GCA		AAA	GTA	AAA					_	•	5374 1663
53		CTT	CGA	GAA		ATT	GCT	AAT	CTG			CTT L						••	CCT	_		5434 1683
54	35	TCT	AAG		ATC	ACA	GAG	CAC	_	AAA	TGC				ATA			-		••	Ť	5494 1703
54		TTT	TTC		•	ATG	CCA	AGG	GGG		_		_	•	CCG	_	•	•	_	ACG		5554 1723
55		TTA	CTA	AAA	crc	AGG	AGG					-							GGC	GGG	ATA	5614 1743
56		AGT	TCA			R CAT H		ACC	GCC		AAA			CTG			••	-				5674
56	75	- ACT	AGA	v GTG V		r TGC C	CAA	AGC	AAC	AAC		_	_				_		M GGC G	G GTC V	R AAG K	1763 5734 1783
57	_	ACT		•	•	TGC	-											-		•		5794 1803
57		ATA	TCA		TCC	AAA K	GGG	GCA	orc	GTT		CTC	CAA	AAG	ACA	•	GGA	_			-	5854 1823
58		GTC	_			ogc G			GCT		TTC		CTA			TIG	-	_	•	-	_	5914 1843
59		17G	•	ATA		GAA		TCC	AGC	GOG				GGC	AGA	_				_	•	5974 1863
	975	GAA				CCT	ACA	AAA	ATA I	ATG M	AGT S	GCA G	ATC	CAG					AAC N	AGA R	GCA A	6034 1883
)35 384		CTG L	ACC T	GAG E	ATG M	GTC V	AAG K	AAG K	ATA I	ACC T	AGC S	ATG M	AAC N	AGG R	GGA G	GAC D	TTC F	AAG K	CAG	ATT I	6094 1903
	095 904		TTG L	GCA A	ACA T	GGG G	GCA A	GGC G	AAA K	ACC T	ACA T	GAA			AAA K		GTT V	ATA I	GAG	GAG E	ATA I	6154 1923
	155 924		AGA R	CAC H	AAG K	AGA R	GTA V			CTT L	ATA I	CCA P		AGG R	GCA A	GCG A	GCA A	GAG E	TCA S	GTC V	TAC Y	6214 1943
	215 944		TAT Y	ATG M	AGA R	TTG L	AAA K	CAC H	CCA P	AGC S	ATC I	TCT S	TTT F	AAC N	CTA L	AGG R	ATA I	GGG G	GAC D	ATG M	AAA K	6274 1963
		GAG E		GAC D	ATG M	GCA A	ACC T	GGG G	ATA I	ACC T	TAT Y	GCA A		TAC Y	GGG	TAC Y	TTC F	TGC	CAA Q	ATG M	CCT P	6334 1983
		CAA Q		AAG K	CTC L	AGA R		GCT A	ATG M	GTA V	GAA E	TAC Y		TAC Y	ATA I	TTC F	TTA L	GAT D	GAA E	TAC Y	CAT H	6394 2003
	395 004		OCC	ACT T	CCT P	GAA E	CAA	CTG L	GCA A	ATT	ATC	GGG	AAG K	ATC I	CAC H	AGA R	TTT F	TCA S	GAG E	AGT S	ATA I	6454 2023
	455 024		GTT V	GTC V	GCC	ATG M		GCC A	ACG T			GGG		GTG V	ACC	ACA T	ACA T	GGT G	CAA Q	AAG K	CAC H	6514 2043
	515 044		ATA I	GAG E	GAA E	TTC F		GCC			GTA V		AAA K		GAG E	GAT D	CTT L	GOT	AGT S	CAG Q	TTC	6574 2063
6 2	575 064	CTT L	GAT D	ATA I	GCA A	GGG	TTA L	AAA K	ATA	CCA P	CTG	GAT D	GAG E	ATG H	AAA K	GGC	AAT N	ATG M	TTG L	GTT V		6634 2083
6	635 084	GTA V	CCA P	ACG T	AGA R	AAC N	ATG M	GCA A	GTA V	GAG E	GTA V	GCA A	AAG K	AAG K	CTA L	AAA K	GCT A	AAG K	GGC G	TAT Y	AAC N	6694 2103
6	695 104	TCT S	OGA G	TAC	TAT Y	TAC Y	AGT S	GGA	GAG E	GAT D	P CCA	GCC A	AAT N	CTG	AGA R	om V	, cua	ACA T	TCA S		TCC	6754 2123
			TAT Y		ATC I	org V	GCT A	ACA T	TAA N	GCT A	I I	GAA E	TCA S	GGA G	OTG V	ACA T	CTA	CCA P	GAT D	TTG	GAC D	6814 2143
6	815 144	ACG T	V V	I I	GAC D	ACC T	G	L L	AAA K	C	GAA E	AAG K	R R	org V	AGC R	CTA V	TCA S	TCA S	AAG K	ATA I	CCC	6874 2163
	875 164		ATC I	ÇT)	T AC	c GCC		K K	AGC R		GCC A	CTC V	ACT T	A CMC	G	GAC E	CAG Q	GCG A	CAG Q	CGT R	AGG R	6934 2183
		GCC		GT/ V	G		V GTG			G		TAT Y			S AGC		GAA E		GCA A	ACA T	, ccc	6994 2203

BVDV NADL (inf. clone) -> Gt 4/21/99 6995 TCA AAG GAC TAC CAC TAT GAC CTC TTG CAG GCA CAA AGA TAC GGG ATT GAG GAT GGA ATC 2204 S K D Y H Y D L L Q A Q R Y G I E D G I 7054 7055 AAC GTG ACG AAA TCC TTT AGG GAG ATG AAT TAC GAT TGG AGC CTA TAC GAG GAG GAC AGC 2224 N V T K S F R E H N Y D W S L Y E E D S 7114 7115 CTA CTA ATA ACC CAG CTG GAA ATA CTA AAT AAT CTA CTC ATC TCA GAA GAC TTG CCA GCC 7174 7175 GCT GTT AAG AAC ATA ATG GCC AGG ACT GAT CAC CCA GAG CCA ATC CAA CTT GCA TAC AAC 7234 2283 7235 AGC TAT GAA GTC CAG GTC CCG GTC CTG TTC CCA AAA ATA AGG AAT GGA GAA GTC ACA GAC 2284 S Y E V O V P V L F P K I R N G E V T D 7294 7295 ACC TAC GAA AAT TAC TCG TTT CTA AAT GCC AGA AAG TTA GGG GAG GAT GTG CCC GTG TAT 2304 T Y E lpha Y S F lpha N A R K lpha G E D V P V Y 7354 2323 7355 ATC TAC OCT ACT GAA GAT GAG GAT CTG GCA GTT GAC CTC TTA GGG CTA GAC TGG CCT GAT 2324 I Y A T E D E D L A V D L L G L D W P D 7414 7415 CCT GGG AAC CAG CAG GTA GTG GAG ACT GGT AAA GCA CTG AAG CAA GTG ACC GGG TTG TCC 2344 P G N Q Q V V E T G K A L K Q V T G L S 7474 2363 7475 TCG GCT GAA AAT GCC CTA CTA GTG GCT TTA TTT GGG TAT GTG GCT TAC CAG GCT CTC TCA 2364 S A E N A L L V A L F G Y V G Y Q A L S 7534 7535 AAG AGG CAT GTC CCA ATG ATA ACA GAC ATA TAT ACC ATG GAG GAC CAG AGA CTA GAA GAC 2384 K R H V P M I T D I Y T I E D Q R L E D 7594 2403 7595 ACC ACC CAC CTC CAG TAT GCA CCC AAC GCC ATA AAA ACC GAT GGG ACA GAG ACT GAA CTG 2404 T T H L Q Y A P N A I K T D G T E T E L 7654 2423 7655 AAA GAA CTG GCG TCG GGT GAC GTG GAA AAA ATC ATG GGA GCC ATT TCA GAT TAT GCA GCT 2424 K E L A S G D V E K I M G A I S D V A A 7714 7715 GGG GGA CTG GAG TIT GTT ANA TCC CAA GCA GAA AAG ATA ANA ACA GCT CCT TTG TTT AAA 2444 G G L E F V K S Q A E K I K T A P L F K 7774 2463 7775 GAA AAC GCA GAA GCC GCA AAA GGG TAT GTC CAA AAA TTC ATT GAC TCA TTA ATT GAA AAT 2464 E N A E A A K G Y V Q K F I D S L I E N 7834 7835 AAA GAA GAA ATA ATC AGA TAT GGT TTG TGG GGA ACA CAC ACA GCA CTA TAC AAA AGC ATA 2484 K E E I I R Y G L W G T H T A L Y K S I 7894 2503 7895 GCT GCA AGA CTG GGG CAT GAA ACA GGG TTT GCC ACA CTA GTG TTA AAG TGG CTA GCT TTT 2504 A A R L G H E T A F A T L V L K W L A F 7954 7955 GGA GGG GAA TCA GTG TCA GAC CAC GTC AAG CAG GCG GCA GTT GAT TTA GTG GTC TAT TAT 2524 G G E S V S D H V K O A A V D L V V Y Y 8014 8015 GTG ATG AAT AAG CCT TCC TTC CCA GGT GAC TCC GAG ACA CAG CAA GAA GGG AGG CGA TTC 2544 V H N K P S F P G D S E T Q Q E G R R F 2563 8075 GTC GCA AGC CTG TTC ATC TCC GCA CTG GCA ACC TAC ACA TAC AAA ACT TGG AAT TAC CAC 2564 V A S L F I S A L A T Y T Y K T W N Y H 8134 8135 AAT CTC TCT AAA GTG GTG GAA CCA GCC CTG GCT TAC CTC CCC TAT GCT ACC AGC GCA TTA 2584 N L S K V V E P A L A Y L P Y A T S A L 8194 2603 8195 AAA ATG TTC ACC CCA ACG CGG CTG GAG AGC GTG GTG ATA CTG AGC ACC ACG ATA TAT AAA 2604 K M F T P T R L E S V V I L S T T I Y K 8254 2623 8255 ACA TAG CTC TCT ATA AGG AAG GAG AAG AGT GAT GGA TTG CTG GGT ACG GAG ATA AGT GCA 2624 T Y L S I R K G K S D G L L G T G I S A 8314 8315 GCC ATG GAA ATC CTG TCA CAA AAC CCA GTA TCG GTA GGT ATA TCT GTG ATG TTG GGG GTA 2644 A M E I L S Q N P V S V G I S V M L G V 8374 8375 GGG GCA ATC GCT GCG CAC AAC GCT ATT GAG TCC AGT GAA CAG AAA AGG ACC CTA CTT ATG 2664 G A I A A H N A I E S S E Q K R T L L M 8434 2683 8435 AAG GTG TTT GTA AAG AAC TTC TTG GAT CAG GCT GCA ACA GAT GAG CTG GTA AAA GAA AAC 2684 K V F V K N F L D O A A T D F L V K F 8494 8495 CCA GAA AAA ATT ATA ATG GCC TTA TTT GAA GCA GTC CAG ACA ATT GGT AAC CCC CTG AGA 2704 P E K I I H A L F E A V Q T I G N P L R 8554 8555 CTA ATA TAC CAC CTG TAT GGG GTT TAC TAC AAA GGT TGG GAG GCC AAG GAA CTA TCT GAG 2724 L 1 Y H L Y G V Y Y Y K G W E A K E L S E 8614 2743

8615 AGG ACA GCA GCC AGA AAC TTA TTC ACA TTG ATA ATG TTT GAA GCC TTC GAG TTA TTA GGG 2744 R T A G R N L F T L I M F E A F E L L G

8675 ATG GAC TCA CAA GGG AAA ATA AGG AAC CTG TCC GGA AAT TAC ATT TTG GAT TTG ATA TAC

FIGURE 11-5

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Page 5

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Page

27/67 BVDV NADL (inf. clone) -> Gc 4/21/99 8735 GGC CTA CAC AAG CAA ATC AAC AGA GGG CTG AAG AAA ATG GTA CTG GGG TGG GCC CCT GCA 2784 G L K K M V L G W A P A 8794 2803 8795 CCC TTT AGT TGT GAC TGG ACC CCT AGT GAC GAG AGG ATC AGA TTG CCA ACA GAC AAC TAT 2804 P F S C D W T P S D E R I R L P T D N Y 8854 8855 TTG AGG GTA GAA ACC AGG TGC CCA TGT GGC TAT GAG ATG AAA GCT TTC AAA AAT GTA GGT 2824 L R V E T R C P C G Y E M K A F K N V G 8914 2843 8915 GGC AAA CTT ACC AAA GTG GAG GAG AGC GGG CCT TTC CTA TGT AGA AAC AGA CCT GGT AGG 2844 G K L T K V E E S G P F L C R N R P G B 8974 8975 GGA CCA GTC AAC TAC AGA GTC ACC AAG TAT TAC GAT GAC AAC CTC AGA GAG ATA AAA CCA 9034 2883 9035 GTA GCA AAG TTG GAA CAG CAG GTA GAG CAC TAC TAC AAA GGG GTC ACA GCA AAA ATT GAC 2884 V A K L E G Q V E H Y Y K G V T A K T GAC 9094 9095 TAC AGT AAA GGA AAA ATG CTC TTG GCC ACT GAC AAG TGG GAG GTG GAA CAT GGT GTC ATA 9154 9155 ACC AGG TTA GCT AAG AGA TAT ACT GGG GTC GGG TTC AAT GGT GCA TAC TTA GGT GAC GAG 2924 T R L A K R Y T G V G F N G A Y L G D F 9214 2943 9215 CCC AAT CAC CGT GCT CTA GTG GAG AGG GAC TGT GCA ACT ATA ACC AAA AAC ACA GTA CAG 2944 P N H R A L V E R D C A T I T K N T V O 9274 9275 TTT CTA AAA ATG AAG AAG GGG TGT GCG TTC ACC TAT GAC CTG ACC ATC TCC AAT CTG ACC 2964 F I. K M K K G C A F T Y D L T I S N L T 9334 2983 9335 AGG CTC ATC GAA CTA GTA CAC AGG AAC AAT CTT GAA GAG AAG GAA ATA CCC ACC GCT ACG 2984 R L I E L V H R N N L E E K E I P T A T 9394 3003 9395 GTC ACC ACA TGG CTA GCT TAC ACC TTC GTG AAT GAA GAC GTA GGG ACT ATA AAA CCA GTA 3004 V T T W L A Y T F V N E D V G T I K P V 9454 9455 CTA GGA GAG AGA GTA ATC CCC GAC CCT GTA GTT GAT ATC AAT TTA CAA CCA GAG GTG CAA 3024 L G E R V I P D P V V D I N L Q P E V O 9515 GTG GAC ACG TCA GAG GTT GGG ATC ACA ATA ATT GGA AGG GAA ACC CTG ATG ACA ACG GGA 3044 V D T S E V G I T I I G R E T L M T T G 9574 3063 9575 GTG ACA CCT GTC TTG GAA AAA GTA GAG CCT GAC GCC AGC GAC AAC CAA AAC TCG GTG AAG 3064 V T P V L E K V E P D A S D N Q N S V K 9634 9635 ATC GGG TTG GAT GAG GGT AAT TAC CCA GGG CCT GGA ATA CAG ACA CAT ACA CTA ACA GAA 3084 I G L D E G N Y P G P G I Q T H T L T E 3103 9695 GAA ATA CAC AAC AGG GAT GCG AGG CCC TTC ATC ATC ATC CTG GGC TCA AGG AAT TCC ATA 3104 E I H N R D A R P F I M I L G S R N S I 9754 9755 TCA AAT AGG GCA AAG ACT GCT AGA AAT ATA AAT CTG TAC ACA GGA AAT GAC CCC AGG GAA 3124 S N R A K T A R N I N L Y T G N D P R E 3147 9815 ATA CGA GAC TTG ATG GCT GCA GGG CGC ATG TTA GTA GTA GCA CTG AGG GAT GTC GAC CCT 3144 I R D L M A A G R M L V V A L R D V D P 9874 9875 GAG CTG TCT GAA ATG GTC GAT TTC AAG GGG ACT TTT TTA GAT AGG GAG GCC CTG GAG GCT 3164 E L S E M V D F K G T F L D R E A L E A 9935 CTA AGT CTC GGG CAA CCT AAA CCG AAG CAG GTT ACC AAG GAA GCT GTT AGG AAT TTG ATA 3184 L S L G Q P K P K Q V T K E A V R N L I 9994 9995 GAA CAG AAA AAA GAT GTG GAG ATC CCT AAC TGG TTT GCA TCA GAT GAC CCA GTA TTT CTG
3204 E Q K K D V E I P N W F A S D D P V F I 10054 3223 10055 GAA GTG GCC TTA AAA AAT GAT AAG TAC TAC TTA GTA GGA GAT GTT GGA GAG CTA AAA GAT 3224 E V A L K N D K Y Y L V G D V G E L K D 10114 10115 CAA GCT AAA GCA CTT GGG GCC ACG GAT CAG ACA AGA ATT ATA AAG GAG GTA GGC TCA AGG 10174 3263 10175 ACG TAT GCC ATG AAG CTA TCT AGC TGG TTC CTC AAG GCA TCA AAC AAA CAG ATG AGT TTA 3264 T Y A M K L S S W F L K A S N K Q M S L 10234 10235 ACT CCA CTG TTT GAG GAA TTG TTG CTA CGG TGC CCA CCT GCA ACT AAG AGC AAT AAG GGG 3284 T P L F E E L L L R C P P A T K S N K G 10294 10295 CAC ATG GCA TCA GCT TAC CAA TTG GCA CAG GGT AAC TGG GAG CCC CTC GGT TGC GGG GTG 3304 H H A S A Y Q L A Q G N W E P L G C G V 10354 10355 CAC CTA GGT ACA ATA CCA GCC AGA AGG GTG AAG ATA CAC CCA TAT GAA GCT TAC CTG AAG 3324 H $^{\circ}$ L $^{\circ}$ G T $^{\circ}$ I $^{\circ}$ P $^{\circ}$ A $^{\circ}$ R $^{\circ}$ V K $^{\circ}$ I H $^{\circ}$ P $^{\circ}$ E $^{\circ}$ A $^{\circ}$ Y $^{\circ}$ L $^{\circ}$ K 3343 10474

BVDV	/ N	IAD	L (i	nf. (clone	·) ·>	- G.	.s			28/6	7									4/	21/99
	75 (64 1							CTT L								AAC N					AAA K	10534 3383
	35 A			AAC N				CTA L			CAG 0									AAC N	ATC I	10594 3403
	95 1		AAC N		CAG O			ACT T		ATG H						AGG R			AAA K	TTG L	CCA P	10654 3423
106		ATA	crc		OCC.	-	ACC	GAC		AAA	ACC		CAT H	GAG E	GCA	ATA	AGA	GAT	AAG	ATA	GAC D	10714 3443
107		AAG	AGT			CCC	CAA	AAT	CCA	GAA	TTG	CAC	AAC	_	TTG	_	GAG		TTC	_	-	10774 3463
107	75	ATA	_	_		•	crg	AAA	CAC	ACC	TAC	CCT	GAG	CTG			GAG		•	••	GCG	10834 3483
108	64 35 84	occ			AGA	AAG	_	GCA	GCA	GGC	TTC		GAG	AAG	AAG		ATC	GGA	GAA		••	10894 3503
108		GAT	_			CAC	CTG	GTA	GAA	CAA	TTG		AGG				GCC	GGG	-	•	ATA I	10954 3523
109		AAA		-				ATA I	CCA			GAG					AGT				•	11014 3543
110	15			GAC				GAG	AAG	AGG	CCA		GIT		CAA		ССТ				-	11074
110		ACG	CTA		_	ACT T		GTC	ATG									GTT		•••		11134 3583
111		GGA	_		GGA	AAG K	ACC	ccc	TTG	TTC						_	AGA	AAG K	GAA E	TGG W	GAC D	11194 3603
	95							GCC A	GTA V		TTT F	GAC D	ACC T	AAA K	GCC		GAC D	ACT T	CAA Q	GTG V	ACT T	11254 3623
	255 524		AAG K	GAT D	CTG L	CAA	CTT L	ATT I	GGA G	GAA E	ATC I	CAG Q	AAA K	TAT Y	TAC Y	TAT Y	AAG K	AAG K	GAG E	TGG W	CAC H	11314 3643
	315 544		TTC F	ATT	GAC D	ACC	ATC I	ACC T	GAC D	CAC H	ATG M	ACA T	GAA E			GTT V		ACA T	GCA A	GAT D	CCT G	11374 3663
	375 564		GTA V		ATA I		AAT N	GGG G		AGA R	GGG G	AGC S	GGC	CAG Q	CCA P	GAC D	ACA T	agt S	GCT A	GCC G	AAC N	11434 3683
	435 684		ATC	TTA L	AAT N	GTC	CTG	ACA T	ATG H	ATG M	TAC Y	GCC	TTC F	TGC	GAA E		ACA T	GGG G	GTA V	CCG P	TAC Y	11494 3703
	495 704		AG1	TTC F		AGG R		GCA A		ATC		GTC V		GGG		GAT D	GCC	TTC F	TTA L	ATA I	ACT T	11554 3723
	555 724		AAJ K		TTA L		r CIG	AAA K	TTT F		AAC N	K K	G G	ATG M	CAG Q	ATT I	CTT	CAT H	GAA E	GCA A	GGC	11614 3743
	615 744		CC7	CAC Q	K AAC	ATA I	ACG T	GAA E	GGG	GAA E	AAG K	ATC M	K	GM V	y GCC	TAT Y	AGA R	TTT	GAG E	GAT D	ATA	11674 3763
	675 764		TTC F	c TGT	TCT S	H H	T ACC	CCA P	GTC V				TOC W	TCC S		N N	ACC T	AGT S	AGT S	H CAC	ATG M	11734 3783
117	735 784	GCC A	G G	R AG	CAC D	ACC T	GCT A	A CLC	I I	CT)	TC) S	K K	ATC M	y S GCN	ACJ T	AGJ R	TTG L	GAT D	TCA S	AGT S	GGA G	11794 3803
								TAT Y													TCC	11854 3823
								R ACC													GAC D	11914 3843
								rat 1 Y			K				A AT		A GCC		r K		GTA V	11974 3863
								r ga													N N	12034 3883
								G													G GAC	12094 3903
		i TC				T GG	G AA K	A GA	A GA	G GO	C AAI N	C TO	G CT	A GT V	T AA N	C GC	C GA(R AG	G CTO	G ATA	A TCC S	12154 3923
		S AG	C AA K					A TA	C AT	A CC	T GA	T AA K	A GG			A TT		A OG.	A AA	G CA	TAT Y	12214 3943

PCT/US99/08850

WO 99/55366

BVDV	NA	\D	L (i	inf.	clon	e) ->	- G	s		:	29/6	7									4/	21/99	5:42:22 PM	Page	8
	5 G/		CAA Q	CTG L		CTA L			GAG E		AAC N		GTC V	ATG M	GGG	CTT V	c ccc	ACT T	GAG E	AGA R	TAC Y	12274 3963			
	5 A			GGT G	CCC P	ATA 1	CTC V	AAT N	CTG L	CTG L	CTG L		AGG R	TTG L		ATT	CTG L				GCC A	12334 3983			
	5 G		GGC G			AGC S	ŤGA	gac	BBBB	tgta	tata	ttgt	aaat	aaat	taat	ccat	gtaca	atag	tgta	tata	aatat	12408 3989			
1240	9 a	gtt	999	accg	tcca	cctc	aaga	agac	gaca	cgcc	caac	acgc	acag	ctaa	acag	tagt	caaga	atta	tcta	cctc	aagat	12488			
1248	9 a	aca	cta	catt	taat	gcac	acag	cact	ttag	ctgt	atga	ggat	acgc	ccga	cgtc	tata	gttg	gact	aggga	aaga	cctct	12568	1		
1250	59 a	aca	gcc	ccc																		12578			

FIGURE 11-8

FIGURE 12-1

30/67

BVDV NADL clns- (inf. clone) -> Genes

DNA sequence 12308 b.p. gtatacgagaat ... ctaacagccccc linear

1	gtat	acga	gaat	taga	aaac	ıgcac	teat	atac	otat	taac	caat	taaa	aata	1818	rtac	iaeci	aaar	1220		cctc	90
																				gtteg	
																				gtgg	
																				gaata gaata	
				gtgo																	394
1			2.					•						,		.994	M	E	L		3
	ATC I	ACA T	AAT N	gaa E	CTT L	TTA L	TAC Y	K K	ACA T	TAC Y	AAA K	CAA Q	AAA K	CCC P	V GTC	GGG	A CLC	GAG E	gaa E	CCT P	454 23
	GTT	TAT	GAT	CAG	GCA	ССТ	GAT	CCC	TTA	TTT	C	GAA	AGG	GGA	GCA	GTC	CAC	CCT	CAA	TCG	514
	V	Y	D	Q	A	С	D	P	L	F	CCT	E	R	G	A	V	H	P	Q	S	43
515	ACG	CTA	aag	CTC	CCA	CAC	aag	aga	GCG	GAA	COC	gat	GTT	CCA	ACC	AAC	TTG	GCA	TCC	TTA	574
44	T	L	K	L	P	H	K	R	G	E	R	D	V	P	T	N	L	A	S	L	63
575	CCA	AAA	AGA	OCT	GAC	TGC	AGG	TCG	GCT	aat	AGC	AGA	GGA	CCT	A	AGC	GCC	ATC	TAC	CTG	634
64	P	K	R	G	D	C	R	S	G	N	S	R	G	P	QLC	S	G	I	Y	L	83
635	aag	CCA	GGG	CCA	CTA	TTT	TAC	CAG	GAC	TAT	AAA	GGT	CCC	GTC	TAT	CAC	AGG	GCC	CCG	CTG	694
84	K	P	G	P	L	F	Y	Q	D	Y	K	G	P	V	Y	H	R	A	P	L	103
695	GAG	CTC	TTT	GAG	GAG	GGA	TCC	ATG	TGT	GAA	ACG	ACT	AAA	CGG	ATA	GGG	AGA	GTA	ACT	GGA	754
104	E	L	F	E	E	G	S	M	C	E	T	T	K	R	I	G	R	V	T	G	123
755	AGT	GAC	GGA	AAG	CTG	TAC	CAC	ATT	TAT	CTC	TGT	ATA	GAT	GGA	TCT	ATA	ATA	ATA	AAA	AGT	814
124	S	D	G	K	L	Y	H	I	Y	V	C	I	D	G	C	I	I	I	K	S	143
815	GCC	ACG	AGA	AGT	TAC	CAA	AGG	GTG	TTC	AGG	TGG	GTC	CAT	AAT	AGG	CTT	GAC	TGC	CCT	CTA	874
144	A	T	R	S	Y	Q	R	V	F	R	W	V	H	N	R	L	D	C	P	L	163
875	TGG	GTC	ACA	ACT	TGC	TCA	GAC	ACG	AAA	GAA	GAG	GGA	GCA	ACA	AAA	AAG	AAA	ACA	CAG	AAA	934
164	W	V	T	T	C	S	D	T	K	E	E	G	A	T	K	K	K	T	0	K	183
935	CCC	GAC	AGA	CTA	GAA	AGG	GGG	AAA	ATG	AAA	ATA	GTC	CCC	AAA	GAA	TCT	GAA	AAA	GAC	AGC	994
184	P	D	R	L	E	R	G	K	N	K	I	V	P	K	E	S	E	K	D	S	203
995	AAA	ACT	AAA	CCT	CCG	GAT	GCT	ACA	ATA	GTG	GTG	GAA	GGA	GTC	AAA	TAC	CAG	GTG	AGG	AAG	1054
204	K	T	K	P	P	D	A	T	I	V	V	E	G	V	K	Y		V	R	K	223
1055	AAG	GGA	AAA	ACC	AAG	agt	AAA	AAC	ACT	CAG	GAC	GGC	TTG	TAC	CAT	AAC	AAA	AAC	AAA	CCT	1114
224	K	G	K	T	K	S	K	N	T	Q	D	G	L	Y	H	N	K	N	K	P	243
1115	CAG	GAA	TCA	CGC	AAG	AAA	CTG	GAA	AAA	GCA	TTG	TTG	GCG	TGG	GCA	ATA	ATA	GCT	ATA	GTT	1174
244	Q	E	S	R	K	K	L	E	K	A	L	L	A	W	A	I	I	A	I	V	263
1175	TTG	TTT	CAA	GTT	ACA	ATG	GGA	GAA	AAC	ATA	ACA	CAG	TOG	AAC	CTA	CAA	GAT	AAT	GGG	ACG	1234
264	L	F	Q	V	T	H	G	E	N	I	T	Q		N	L	Q	D	N	G	T	283
1235	GAA	GGC	ATA	CAA	CGG	GCA	ATG	TTC	CAA	AGG	GOT	GTG	aat	AGA	ACT	TTA	САТ	GGA	ATC	TGG	1294
284	E	G	I	Q	R	A	M	F	Q	R	G	V	N	R	S	L	Н	G	I	W	303
1295 304	CCA P	GAG E	AAA K	ATC	TCT C	ACT T	COT	GTC V	CCT P	TCC S	CAT H	CTA L	GCC A	ACC T	GAT D	ATA I	GAA E	CTA L	AAA K	ACA T	1354 323
1355 324		CAT H	CCT C	ATG M	ATG M	GAT D	GCA A	AGT S	GAG E	AAG K	ACC T	AAC N	TAC Y	ACG T	TCT C	TGC C	AGA R	CTT L	CAA	CGC R	1414 343
1415		GAG	TGG	AAC	AAG	CAT	GGT	TCG	TCC	AAC	TGG	TAC	AAT	ATT	GAA	CCC	TGG	ATT	CTA	GTC	1474
344		E	W	N	K	H	G	W	C	N	W	Y	N	I	E	P	W	I	L	V	363
1475	ATG	AAT	AGA	ACC	CAA	GCC	AAT	CTC	ACT	GAG	GGA	CAA	CCA	CCA	AGG	GAG	TGC	GCA	GTC	ACT	1534
364	H	N	R	T	Q		N	L	T	E	G	Q	P	P	R	E	C	A	V	T	383
1535 384	TGT C	AGG R	TAT Y	GAT D	AGG R	GCT A	AGT S	GAC D	TTA L	AAC N	CTC V	GTA V	ACA T						CCC P	ACA T	1594 403
1595	CCC	TTA	ACA	GCT	100	aag	AAA	GGA	AAG	AAC	TTC	TCC	TTT	GCA	GGC	ATA	TTG	ATG	CGG	GGC	1654
404		L	T	G	C	K	K	G	K	N	F	S	P	A	G	I	L	M	R	G	423
1655 424	CCC P	TGC C	AAC N	TTT F	GAA E	ATA I	GCT A	GCA A	AGT S	GAT D	GTA V	TTA L	TTC F	AAA K	GAA E	CAT H	GAA E	CGC R			1714 443
1715 444			CAG Q	GAT D	ACT T	ACT T	CTT L	TAC Y	CTT L	GTT V	GAC D	GGG G	TTG L	ACC T	AAC N	TCC S	TTA L	GAA E	_	QCC	1774 463

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BVDV	N A D	, al	 -	lint	clos	ne)	(Sene		31/6	7									4/	21/99
1775		CAA	GGA	ACC	ост	AAA	CTG	ACA	ACC	TGG	TTA L	GGC	AAG K	CAG	crc L	ccc G	ATA I	CTA L			1834 483
1835 484	AAG	TTG	GAA	AAC	AAG	AGT	AAG	ACG		TTT	GGA	GCA	TAC	CCT			CCT		TGT	GAT D	1894 503
1895 504	GTC	GAT	ccc	 AAA	ATT	•	•			TAT	ACA	AAA				-	GCC		TTA	CCC	1954 523
1955 524	AAG	AAC	ACA		ATT		GGC	сст		AAA K	TT									ATA I	2014 543
2015 544	TTA	CAT					CAC H	TTG L				CTA L							CTG L	TCC S	2074 563
2075 564	GAC D					ACA T	GCT A	AGT S	GTA V	ATG M			ATC		CAT H	TTT F	TCC S	ATC I	CCA P	CAA O	2134 583
2135 584	ACT S					ATG M	GAT D					CAG Q		AAC N			GTG V	GAG E	CTG L	ACA T	2194 603
2195 604	ACA T	GCT A					GGG G		GTC V		AAT N				TAT Y			ATA I		CCA P	2254 623
2255 624	AAT N		TGG W	CCT P		GAG E	ACA T	ACT T	GTA V		TTG L	GCA A	TTT F	GAA E	GAG E	GTG V		CAG Q	GTG V	GTG V	2314 643
2315 644	AAG K	TTA L	GTG V				CTC L		GAT D	TTA L	ACA T	CGC R	ATT I	TGG W	AAC N	GCT A		ACA T	ACT T	ACT T	2374 663
2375 664	GCT A	TTT F		GTA V	TGC C	CTT L	GTT V	aag K	ATA I	GTC V	AGG R	G G G		ATG H	GTA V	CAG Q	GGC G	ATT I	CTG C	TGG W	2434 683
2435 684	CTA L	CTA L	TTG L	ATA I			GTA V	CAA Q			TTG L				CCT P	gaa E	TTC F	TCG S	TAT Y	OCC A	2494 703
2495 704	ATA I	GCA A		GAC D	gaa e		ATT I							GGC G				ACT T	TGG W	aag K	2554 723
2555 724	GAA	TAC Y	TCA S	CCT P	GGA G	ATG M	AAG K	CTG L	gaa e	GAC D	ACA T	ATG M	orc orc	ATT I	GCT A	TGG W	TGC C	gaa E	GAT D	GCG G	2614 743
	AAG K	TTA L	ATG M	TAC Y	CTC	CAA Q	AGA R	TGC C	ACG T	aga R	GAA E	ACC T	AGG R	TAT Y	CTC L	GCA A	ATC I	r TTG	CAT H	ACA T	2674 763
	AGA R	GCC A	TTG L	P CCC	ACC T		v GTG			AAA K				GAT D	GCC G	CGA R	AAG K	CAA Q	GAG E	GAT D	2734 783
	GTA V	GTC V	gaa E	ATG M	AAC N	GAC D	AAC N	TTT P	GAA E	TTT F		CTC L	TGC C	CCA P	TGT C	GAT D	GCC A	AAA K	CCC P	ATA I	2794 803
	GTA V	AGA R	GGG	AAG K	TTC F	AAT N	ACA T			CTG L				GCC A	TTC F	CAG Q	ATG M	GTA V	TGC C	P CCC	2854 623
	ATA I		TGG W	ACA T	G G	ACT T	GTA V	AGC S	C	T ACG		TTC F	AAT N	ATG M	GAC D	ACC T	TTA L	GCC A	ACA T	ACT T	2914 843
	ong V	GTA V	CGG R	ACA T	TAT Y	aga R	AGG R	TCT S	K K	P CCA	F	P	H	AGG R	Q Q	c	TGT C	ATC I	ACC T	CAA Q	2974 863
864	AAC K	N	L	G	E	D	L	н	N	С	I	L	G	G	N	W	T	С	٧	P	3034 883
884	G	D	Q	L	L	Y	ĸ	G	G	S	I	E	S	С	K	W	С	G	Y	0	3094 903
904	F	ĸ	E	S	Ε	G	L	P	H	Y	P	1	G	K	С	ĸ	L	E	N	GAG E	923
92	ACT	G	Y	R	L	٧	D	S	Т	S	С	N	R	E	G	ν	A	I	V	₽	3214 943
94	S CAU	G	т	L	K	С	K	I	G	K	T	T	V	Q	V	ı	A	M	D	T	3274 963 3334
96	K	L	G	P	M	P	c	R	P	Y	E	I	I	s	s	E	G	P	V	E	983
98	4 K	T	λ	С	T	F	N	Y	T	K	T	L	K	N	K	Y	P	E	P	R	1003
100	4 D	s	Y	F	0	o CCC	Y	М	L TT	K GCT	G GAC	E	Y	Q	Y	W	F	D	L	Ε	1023 3514
		T	D	н	н	R	D	Y	F	A	Ε	S	1	L	V	V	ν	V		L	1043

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32/67 BVDV NADL cins- (inf. clone) Genes 4/21/99 3515 TTG GGT GGC AGA TAT GTA CTT TGG TTA CTG GTT ACA TAC ATG GTC TTA TCA GAA CAG AAG 3575 GCC TTA GGG ATT CAG TAT GGA TCA GGG GAA GTG GTG ATG ATG GGC AAC TTG CTA ACC CAT 1064 A L G I Q Y G S G E V V M N G N L L T H 3635 AAC AAT ATT GAA GTG GTG ACA TAC TTC TTG CTG CTG TAC CTG CTG CTG AGG GAG GAG GAG LORA N N T E V V T Y E L L Y L L R E E S 3694 1103 3695 GTA AAG AAG TGG GTC TTA CTC TTA TAC CAC ATC TTA GTG GTA CAC CCA ATC AAA TCT GTA
1104 V K K W V L L L Y H I L V V H P I K S V 3754 1123 3755 ATT CTG ATC CTA CTG ATG ATT GOG GAT GTG GTA AAG GCC GAT TCA GOG GCC CAA GAG TAC 3814 3815 TTG GGG AAA ATA GAC CTC TGT TTT ACA ACA GTA GTA CTA ATC GTC ATA GGT TTA ATC ATA 3874 1163 3875 GCC AGG CGT GAC CCA ACT ATA GTG CCA CTG GTA ACA ATA ATG GCA GCA CTG AGG GTC ACT 1164 A R R D P T I V P L V T I M A A L R V T 3934 1183 3935 GAA CTG ACC CAC CAC CGG GGT GGT GAC ATC GCT GGG GCG GTC ATG ACT ATA ACC CTA CTG 1184 E L T H Q P G V D I A V A V M T I T L L 3994 1203 3995 ATG GTT AGC TAT GTG ACA GAT TAT TTT AGA TAT AAA AAA TGG TTA CAG TGC ATT CTC AGC 1204 M V S Y V T D Y F R Y K K W L Q C I L S 4054 1223 4055 CTG GTA TCT GCG GTG TTC TTG ATA AGA AGC CTA ATA TAC CTA GGT AGA ATC GAG ATG CCA 4114 4115 GAG GTA ACT ATC CCA AAC TGG AGA CCA CTA ACT TTA ATA CTA TTA TAT TTG ATC TCA ACA 1244 E V T I P N W R P L T L I L L Y L I S T 1263 4175 ACA ATT GTA ACG AGG TGG AAG GTT GAC GTG GCT GGC CTA TTG TTG CAA TGT GTG CCT ATC 1264 T 1 V T R W K V D V A G L L L O C V P I 4234 4235 TTA TTG CTG GTC ACA ACC TTG TGG GCC GAC TTC TTA ACC CTA ATA CTG ATC CTG CCT ACC 1284 L L L V T T L W A D F L T L I L P T 4294 1303 4295 TAT GAA TTG GTT AAA TTA TAC TAT CTG AAA ACT GTT AGG ACT GAT ATA GAA AGA AGT TGG 4354 4355 CTA GGG GGG ATA GAC TAT ACA AGA GTT GAC TCC ATC TAC GAC GTT GAT GAG AGT GGA GAG 1324 L G G I D Y T R V D S I Y D V D E S G E 4414 1343 4415 GGC GTA TAT CTT TTT CCA TCA AGG CAG AAA GCA CAG GGG AAT TTT TCT ATA CTC TTG CCC 1344 G V Y L F P S R Q K A Q G N F S I L L P 1363 4475 CTT ATC AAA GCA ACA CTG ATA AGT TGC GTC AGC AGT AAA TGG CAG CTA ATA TAC ATG AGT 1364 L 1 K A T L I S C V S S K W O L I Y M S 4534 4535 TAC TTA ACT TTG GAC TTT ATG TAC TAC ATG CAC AGG AAA GTT ATA GAA GAG ATC TCA GGA 1384 Y L T L D F M Y Y M H R K V I E E I S G 4594 1403 4595 GGT ACC AAC ATA ATA TCC AGG TTA GTG GCA GCA CTC ATA GAG CTG AAC TGG TCC ATG GAA 1404 G T N I I S R L V A A L I E L N W S M F 4654 4655 GAA GAG GAG AGC AAA GGC TTA AAG AAG TTT TAT CTA TTG TCT GGA AGG TTG AGA AAC CTA 1424 E E E S K G L K K F Y L L S G R L R N L 1443 4715 ATA ATA AAA CAT AAG GTA AGG AAT GAG ACC GTG GCT TCT TGG TAC GGG GAG GAA GTC 1444 I I K H K V R N E T V A S W Y G E E E V 4774 1463 4775 TAC GOT ATG CCA AAG ATC ATG ACT ATA ATC AAG GCC AGT ACA CTG AGT AAG AGC AGG CAC 1464 Y G M P K I M T I I K A S T L S K S R H 1483 4835 TOC ATA ATA TOC ACT GTA TGT GAG GGC CGA GAG TGG AAA GGT GGC ACC TGC CCA AAA TGT 1484 C ! I C T V C E G R E W K G G T C P K C 4894 4895 GGA COC CAT GGG AAG CCG ATA ACG TOT GGG ATG TCG CTA GCA GAT TTT GAA GAA AGA CAC 1504 G R H G K P I T C G M S L A D F E E R H 4954 1523 4955 TAT AAA AGA ATC TTT ATA AGG GAA GGC AAC TTT GAG GGGGGGCC TTC AGG CAG GAA TAC AAT 5014 5015 GGC TTT GTA CAA TAT ACC GCT AGG GGG CAA CTA TTT CTG AGA AAC TTG CCC GTA CTG GCA 1542 G F V Q Y T A R G Q L F L R N L P V L A 5074 1561 5075 ACT ANA GTA ANA ATG CTC ATG GTA GGC ANC CTT GGA GAN GAN ATT GGT ANT CTG GAN CAT 1562 T K V K M L M V G N L G E E I G N L E H 5135 CTT GGG TGG ATC CTA AGG GGG CCT GCC GTG TGT AAG AAG ATC ACA GAG CAC GAA AAA TGC 1582 L G W I L R G P A V C K K I T E H E K C 5194 1601 5195 CAC ATT AAT ATA CTG GAT AAA CTA ACC GCA TTT TTC GGG ATC ATG CCA AGG GGG ACT ACA N I L D K LT G М G

		_						C		33	/67									4.0	11/04)
BVDV	NAD							Gene		ACC.	TTA	CTA.	AAA	GTG.	AGG	ACC	CCT	CTG	GAG		21/99 5314
1622	P	R	A	P	V	R	F	P	T	s	L	L	K	٧	R	R	G	L	Ē	т	1641
5315 1642	y CCC	TGG W	GCT A	TAC Y	ACA T	CAC H	CAA Q		C			TCA S	GTC V	GAC D	CAT H	GTA V	ACC T	A A	GGA G	AAA K	5374 1661
5375 1662	GAT D	CTA L	CTG L	CTC V	TCT C	GAC D	AGC S					AGA R	otc v	CTT V	TGC C	CAA Q	AGC S	AAC N	AAC N	AGG R	5434 1681
	TTG L	ACC T	GAT D		ACA T	gag E	TAT Y	GGC	GTC V	AAG K	ACT T	GAC D	TCA S	G G	TGC C	CCA P	GAC D	GGT G	GCC A	aga R	5494 1701
5495 1702	TCT C		GTG V	TTA L	AAT N	CCA P	gag E		GTT V		ATA I	TCA S	GGA G	TCC S	AAA K	G G	GCA A	GTC V	য়েয় V	CAC H	5554 1721
5555 1722	CTC L	CAA Q	AAG K	ACA T	GGT G		gaa E	TTC F	ACG T	TGT C	GTC V	ACC T	GCA À	TCA S	GCC	ACA T	CCG P	OCT A	TTC F	TTC F	5614 1741
5615 1742	GAC D	CTA L	AAA K	AAC N		AAA K		TGG W							GAA E	GCC A	TCC S	AGC S	GGG G	agg R	5674 1761
5675 1762	GTG V	GTT V	GGC G	AGA R	GTC V	AAA K	GTA V		AAG K			GAG E	TCT S	AAA K	CCT P	ACA T	AAA K	ATA I	atg M	act S	5734 1781
5735 1782	GGA G	ATC I	CAG Q	ACC T	GTC V	TCA S	AAA K					CTG L	ACC T		ATG M		aag K	AAG K	ATA I	ACC T	5794 1801
5795 1802	AGC S	ATG M	AAC N	AGG R			TTC F	AAG K						ACA T	œc G	GCA A	GGC G	AAA K	ACC T	ACA T	5854 1821
5855 1822	GAA E	CTC L	CCA P	AAA K	GCA A	GTT V	ATA I		GAG E	ATA I	GGA O	AGA R	CAC H	aag K	aga R	GTA V	TTA L	GTT V	CTT L	ATA I	5914 1841
5915 1842	CCA P		AGG R	GCA A	GCG A		GAG E						ATG M		TTG L	K K	CAC H	CCA P	AGC S	ATC I	5974 1861
5975 1862	TCT S	TTT F	AAC N	CTA L	AGG R	ATA I					GAG E	GGG G	GAC D	ATG M	ŒA Å	ACC T	GGC G	ATA I	ACC T	TAT Y	6034 1881
6035 1882	GCA A	TCA S	TAC Y	GGG G	TAC Y	TTC F	TGC C	ÇAA Q	ATG M	CCT P	CAA Q	CCA P	AAG K	CTC L	AGA R	GCT A	GCT A	ATG M	gta V	GAA E	6094 1901
6095 1902	TAC Y	TCA S	TAC Y	ATA I	TTC F	TTA L		GAA E		CAT H	TGT C	GCC A	ACT T	CCT P	GAA E	CAA Q	CTG L	y Cy	ATT I	ATC I	6154 1921
6155 1922	GOG	AAG K	ATC I	CAC H	AGA R	TTT F	TCA S		AGT S		AGG R	GTT V	GTC V	GCC	atg M	ACT T	A GCC	ACG T	CCA P	GCA A	6214 1941
6215 1942	GGG		GTG V	ACC T	ACA T	ACA T			AAG K	CAC H	CCA P	ATA I	GAG E		TTC F	ATA I	GCC A	CCC P	GAG E	GTA V	6274 1961
6275 1962	ATG M	AAA K	GGG G	GAG E					CAG Q			GAT D				TTA L	AAA K	ATA I	CCA P	A QIC	6334 1981
6335 1982	GAT D	GAG E	ATG M	AAA K	GCC G	AAT N	ATG M	TTG L	chi Chi	TTT F	GTA V	CCA P	ACG T	AGA R	AAC N	atg M	GCA A	gta V	gag E	GTA V	6394 2001
6395 2002	GCA A		AAG K	CTA L	AAA K	GCT A	AAG K		TAT Y	AAC N	TCT S	GGA G	TAC Y	TAT Y	TAC Y	AGT S	GGA G	gag E	GAT D	CCA P	6454 2021
6455 2022	A OCC		CTG L	AGA R		GTG V		TCA S					GTA V	ATC I		GCT A	ACA T	AAT N	GCT Å	ATT I	6514 2041
6515 2042	GAA E	TCA S	GGA G	GTG V	ACA T		CCA P					GPT V			ACG T		TTG L	AAA K		gaa E	6574 2061
6575 2062	aag K	AGG R	GTG V	AGG R	GTA V	TCA S	TCA S	AAG K	ATA I	CCC P	TTC F	ATC I	GTA V	ACA T	GGC	CTT L	AAG K	AGG R	ATG M	GCC A	6634 2081
6635 2082	GTG V		ങ v										GTA V					CCC P		AGG R	6694 2101
	TAT Y			AGC S		GAA E		GCA A	ACA T				GAC D						TTG L	CAG Q	6754 2121
	GCA A															TTT F				aat N	6814 2141
6815 2142	TAC Y	GAT D	TGG W	AGC S	CTA L	TAC Y	GAG E	GAG E	GAC D	AGC S	CTA L	CTA L	ATA I	ACC T	CAG Q	CTG L	gaa E	ATA	CTA L	aat N	6874 2161
	AAT N		CTC L				GAC D		CCA P						ATA I	ATG M		AGG R	ACT T	GAT D	6934 2181
6935 2182	CAC H								TAC Y				GAA E		CAG Q			GTC V	CTG L	TTC F	6994 2201

BVDV	NAI	OL c	ins-	(inf	ī. ci	one)		Gen	es		34	1/67								4	/21/99
	CCA			AGG					ACA T	GAC			GAA E			TCC	TTT F	CTA L	AAT	GCC	7054 2221
	AGA		_	•	•		_	-	•	_								_	•-	CA A	7114 2241
7115 2242	GTT				GCC			TOG	CCT		-	GGG		CAG		_			-		7174 2261
7175 2262	AAA	-	_		CAA							GCT	-	Ĭ.	_		•		GCT		7234
7235 2282	TTT		TAT		-		CAG	_	_	_	_			••			ATA I	ACA T	••	_	7294 2301
7295 2302	TAT		-	•	GAC				GAA			ACC		CTC		•-	-	•	_	GCC	7354 2321
7355 2322	ATA	*	-	_		_			GAA	-				_	_			•	•-	A AAA K	7414 2341
7415 2342	ATC		-			TCA										-		•	_		7474 2361
7475 2362	GAA				ACA	GCT	-			AAA	GAA	AAC		GAA	-	•	•		_		7534 2381
7535 2382	CAA	***				TCA	-		-	AAT		GAA						-		•	7594 2401
7595 2402	GGA	ACA						٠.									-		_	TTT	7654 2421
7655 2422					TTA		TGG	CTA		TTT		GGG			_		GAC D	-	GTC V	AAG K	7714 2441
7715 2442		GCG A	GCA A	OLL OLL	GAT D	TTA L	GTG V	GTC V	TAT Y	TAT Y	GTG V				CCT P	TCC	TTC F	CCA P	OGT G	GAC D	7774 2461
7775 2462		GAG E	ACA T	CAG Q	CAA	GAA E	GGG G			TTC F	GTC C	GCA A	AGC S	CTG	TTC F	ATC I	TCC	GCA A	CTG L	GCA A	7834 2481
7835 2482		TAC Y	ACA T	TAC Y	AAA K	ACT T		AAT N	TAC Y		AAT N	CTC L			GTG V	ong V	GAA E	CCA P	GCC A	CTG L	7894 2501
7895 2502		TAC Y	CTC L	CCC P	TAT Y	GCT A	ACC T	AGC S	GCA A		AAA K	ATG M	TTC F	ACC T	CCA P	ACG T	CGG R	CTG L	GAG E	AGC S	7954 2521
7955 2522		GTG V	ATA I	CTG L	AGC S	ACC T	ACG T	ATA I	TAT Y	AAA K	ACA T	TAC Y	CTC L	TCT S	ATA I	AGG R	AAG K	GGG G	AAG K	AGT S	8014 2541
8015 2542				CTG L	GCT G	ACG T	GGG G	ATA I	AGT S		OCC A			ATC I	CTG L	TCA S		AAC N	CCA P	GTA V	8074 2561
8075 2562		GTA V	OGT G	ATA I	TCT S	GTG V	ATG M		GGG G	GTA V	GG G	GCA A	ATC I	GCT A	GCG A	CAC H	AAC N	GCT A	ATT I	GAG E	8134 2581
8135 2582		AGT S			AAA K	AGG R					AAG K		TTT F	GTA V	AAG K	AAC N	TTC F	TTG L	GAT D	CAG Q	8194 2601
8195 2602	GCT A	GCA A	ACA T	GAT D	GAG E	CTG L	GTA V			AAC N			AAA K	ATT I	ATA I	ATG M	GCC A	TTA L	TTT P	GAA E	8254 2621
8255 2622				ACA T	ATT I	ост С							TAC Y		CTG L	TAT Y	occ G	CTT V	TAC Y	TAC Y	8314 2641
8315 2642	AAA K	GGT G	TGG W	gag E	GCC A	AAG K	GAA E	CTA L	TCT S	GAG E	AGG R	ACA T	GCA A	GOC G	AGA R	AAC N	TTA L	TTC F	ACA T	TTG L	8374 2661
8375 2662	ATA I	ATG M	TTT F	gaa E	GCC A	TTC F	GAG E	TTA L	TTA L	GGG G	ATG M	GAC D	TCA S	CAA Q	G G	AAA K	ATA I	AGG R	AAC N	CTG L	8434 2681
8435 2682	TCC S	GGA G	AAT N	TAC Y	ATT I	TTG L	GAT D	TTG L	ATA I	TAC Y	GGC G	CTA L	CAC H	AAG K	CAA Q		AAC N			CTG L	8494 2701
8495 2702	AAG K	AAA K	ATG H	GTA V	CTG L	GGC G	TOG W	GCC A	CCT P	GCA A	CCC P	TTT F	act S	TGT C	GAC D	TGG W	ACC T	CCT P	AGT S	GAC D	8554 2721
8555 2722	GAG E	AGG R	ATC I	aga R	TTG L	CCA P	ACA T	GAC D	AAC N	TAT Y	TTG L	AGG R	CTA V	GAA E	ACC T	AGG R	TGC C	CCA P	TGT C	GCC G	8614 2741
8615 2742																				GGG G	8674 2761
8675 2762	CCT P	TTC F	CTA L	TGT C	aga R	AAC N	aga R	CCT P	G G	agg R	GGA G	CCA P	GTC V	aac N	TAC Y	aga R	GTC V	ACC T	AAG K	TAT Y	8734 2781

PUDU MADI ALIA	(i=f =1===)	Canas	35/67	4/31/00
BVDV NADL cins- 8735 TAC GAT GAC	AAC CTC AG	GAG ATA AAA (CCA GTA GCA AAG TTG GAA GG	4/21/99 A CAG GTA GAG CAC 8794
2782 Y D D	N L R	EIK	PVAKLEG GACTACAGT AAA GGA AAA ATK	Q V E H 2801
2803 A A K	G V T	AKII	у з к с к м	L L A T 2821
2822 D K W	E V E	H G V		Y T G V 2841
8915 GGG TTC AAT 2842 G F N	GCT GCA TAC	L G D E	EAG CCC AAT CAC CGT GCT CT/ E P N H R A L	V E R D 2861
8975 TGT GCA ACT 2862 C A T	ATA ACC AAV		CAG TTT CTA AAA ATG AAG AAC) F L K M K K	G C A F 2881
9035 ACC TAT GAC 2882 T Y D	CTG ACC ATC	TCC AAT CTG A	ACC AGG CTC ATC GAA CTA GT/	CAC AGG AAC AAT 9094 H R N N 2901
9095 CTT GAA GAG 2902 L E E	AAG GAA ATA	CCC ACC GCT /	ACG GTC ACC ACA TGG CTA GCT	TAC ACC TTC GTG 9154 Y T F V 2921
9155 AAT GAA GAC 2922 N E D			TTA CTA GGA GAG AGA GTA ATO	CCC GAC CCT GTA 9214 P D P V 2941
9215 GTT GAT ATC 2942 V D I	AAT TTA CAU	CCA GAG GTG C	CAA GTG GAC ACG TCA GAG GTM	CGG ATC ACA ATA 9274 G I T I 2961
9275 ATT OGA AGG 2962 I G R	GAA ACC CTC		GA GTG ACA CCT GTC TTG GAU	A AAA GTA GAG CCT 9334 K V E P 2981
	•		NG ATC GOG TTG GAT GAG GG	
9395 CCT GGA ATA			SAA GAA ATA CAC AAC AGG GA1	
	•		ATA TCA AAT AGG GCA AAG ACT	
			SAA ATA CGA GAC TTG ATG GCT	
		GAT GTC GAC C	TOT GAS CTG TOT GAA ATG STO	
9635 ACT TTT TTA	GAT AGG GAG	GCC CTG GAG G	CT CTA AGT CTC GGG CAA CCT	ANA CCG ANG CAG 9694
			ITA GAA CAG AAA AAA GAT GTO	
			TG GAA GTG GCC TTA AAA AAT	
			, e v a l k n Pat caa gct aaa gca ctt ggo	D K Y Y 3141 GCC ACG GAT CAG 9874
3142 L V G 9875 ACA AGA ATT	D V G	E L K D	O Q A K A L G OG ACG TAT GCC ATG AAG CTA	A T D Q 3161 TCT AGC TGG TTC 9934
3162 T R I 9935 CTC AAG GCA	I K E	V G S R	TYAMKL TAACT CCACTG TTT GAG GAJ	S S W F 3181
3182 L K A	S N K	Q M S L		L L L R 3201
3202 C P P	ат к	SNKG	HMASAY	Q L A Q 3221
3222 G N W	Ë P L	G C G V	TO CAC CTA GGT ACA ATA CCA H L G T I P	A R R V 3241
10115 AAG ATA CAC 3242 K I H	P Y E	A Y L K	AG TTG AAA GAT TTC ATA GAP	
10175 AAA CCT AGG 3262 K P R	GTT AAG GAT V K D	ACA GTA ATA A T V I R	GA GAG CAC AAC AAA TOG AT# E H N K W I	CTT AAA AAA ATA 10234 L K K I 3281
10235 AGG TTT CAA 3282 R F Q	GGA AAC CTC G N L	AAC ACC AAG A N T K K	AA ATG CTC AAC CCG GGG AAA M L N P G K	CTA TCT GAA CAG 10294 L S E Q 3301
10295 TTG GAC AGG 3302 L D R	GAG OGG CGC E G R	AAG AGG AAC A	TC TAC AAC CAC CAG ATT GGT Y N H Q I G	
			CA ATA GTG AGG GCC CAA ACC	GAC ACC AAA ACC 10414 D T K T 3341
	GCA ATA AGA A I R	CAT AAG ATA G	AC AAG AGT GAA AAC CGG CAA	AAT CCA GAA TTG 10474 N P E L 3361

BVDV	NA	n i	elne.	(in	f d	one)		Gen	20		36/6	67									
		C AA				GAC				: ACC	S AT	A GCC	CA	A CCC	: ACC	: cre	S AA	A CAC	: ACC	TAC	10534
330.	2 H	N	K	L	L	£	1	F	н	T	I	A	Q	P	т	L	К	н	T	Y	3381
338	2 G	ε	V	т	W	E	Q	L	E	A	G	I	N	R	K	G	A	A	G	F TTC	10594 3401
10599 340	S CTO	G GAC	K K	AAG K	N N	I ATC	C C	GAA E	CTA V	L	GAT D	TC/ S	GA/ E	A AAC	CAC H	r CTC	GT/ V	GAJ E	CAA Q	TTG L	10654 3421
10659 3422	GTC	AGC R	GAT D	CTG L	AAC K	GCC A	GCG	AGA R	AAG K	ATA I	AAJ K	TAT Y	TA1	GA/	ACT	GC/	ATA	CCA P	AAA K	AAT N	10714 3441
10719 3442	GAC	AAC K	AGA R	GAT D	orc	AGT S	GAT	GAC D	TGG	CAG	GC/	GGG	GAC D	cro L	GTG V	GM V	GAC	AAG K	AGG	CCA P	10774 3461
10775 3462	AG/	GM V	ATC	CAA	TAC	CCT P	GAA E	GCC	AAG K	ACA	AGG	CTA L	GCC	ATC	ACT	AAG K	GTC	ATG	TAT	AAC N	10834 3481
10835 3482	TGC	e carc	AAA K	CAG	CAG	eccc P	GTT V	GTG V	ATT	CCA P				GGA	_		CCC		-	•	10894 3501
10895	ATC	TTT	GAT D	-	_	AGA R	AAG K				_					-	-		•	••	10954 3521
10955 3522	GAC	ACC	AAA K	GCC A	TCC W	GAC											•	-	-	•	11014 3541
11015 3542	CAG	AAA K	TAT		TAT		AAG K	_			_		_		_		_	-	_		11074
11075 3562	ACA		GTA V									-	_	_	-		-	-		•••	3561 11134 3581
11135 3582	AGC S	occ	CAG	CCA P	GAC		AGT	CCT	GGC				-			•	_	_	••		11194 3601
11195 3602	GCC	TTC F	TGC	GAA E	AGC S	ACA T	GGG G	GTA V				AGT S	TTC F			_	-		••	-	11254 3621
11255 3622	GTC V	TGT C	GOG G	GAT D	GAT D	GGC G	TTC F	TTA L	ATA I	ACT T	GAA E					CIG			_		11314 3641
11315 3642	AAA K	GCG G	ATG M	CAG O	ATT I	CTT L	CAT H	GAA E	GCA A	GGC	AAA K					ACG			••		11374 3661
11375 3662	ATG M	AAA K	GÎT V	GCC A	TAT	AGA R	TTT F	GAG E	GAT D	ATA I	GAG E	TTC F	TGT C	TCT S				_	CCT	GTT V	11434 3681
11435 3682	AGG R	TGG W	TCC S	GAC D	AAC N	ACC T	AGT S	AGT S	CAC H	ATG M				-			_		-	•	11494 3701
11495 3702	AAG K	ATG M	GCA A	ACA T	AGA R	TTG L	GAT D	TCA S	AGT S	GGA G	GAG E	AGG R	GGT G	ACC T	ACA T	GCA A	TAT Y	GAA E	_	GCG	11554 3721
11555 3722	GTA V	GCC A	TTC F	AGT S	TTC F	TTC L	CTG L	ATG M	TAT Y	TCC S	TOG W	AAC N	CCG P	CTT L	GTT V	AGG R	AGG R	ATT I		CTG L	11614 3741
11615 3742	TTG L	GTC V	CTT L	TCG S	CAA Q	CAG Q	CCA P	GAG E	ACA T	GAC D	CCA P	TCA S	AAA K	CAT H	GCC	ACT T	TAT Y	TAT Y			11674 3761
11675 3762	ост G	GAT D	CCA P	ATA I	GGG G	OCC	TAT Y	AAA K	GAT D	GTA V	ATA I			AAT N	CTA L	AGT S	GAA E	CTG L		AGA R	11734 3781
11735 3782	ACA T	ogc G	TTT F	GAG E	AAA K	TTG L	GCA A	aat N	CTA L	AAC N	CTA L	AGC S	CTG L	TCC S	ACG T	TTG L	GGG			ACT T	11794 3801
11795 3802	aag K	CAC H	ACA T	AGC S	AAA K	AGA R	ATA I	ATT I	CAG 0	GAC D	TGT C	GTT V	GCC	ATT I	GGG G	AAA K		GAG		AAC N	11854 3821
11855 3822	TGG W	CTA L	GTT V	AAC N	GCC A	GAC D	AGG R	CTG L	ATA I	TCC S	AGC S	AAA K	ACT T							GAT D	11914 3841
11915 3842	AAA K	c G	TTT F	ACA T	TTA L	CAA Q	GGA G	AAG K	CAT H	TAT Y	GAG E									AAC N	11974 3861
11975 3862	CCG P	GTC V	ATG	GGC	GTT V	GGG G	ACT T	GAG E	AGA R	TAC Y										CTG L	12034 3881
12035 3882	AGA R	AGG R	TTG	AAA . K	ATT I	CTG L	CTC L	ATG	ACG T	GCC A	GTC V	GGC G	GTC V	AGC S	AGC S						
12099	tgta	aata	aatt	aatc	catg	taca	tagt	gtat	асаа	atat	agtt	9998	ccgt	CCAC	ctca	agaa	gacg	acac	gccc	aaca	
12179	cgca	cago	Caaa	cagt	agtc	aaga	ttat	CEAC	ctca	agat	aaca	ctac	attt	aatg	caca	cagc	actt	tage	tgta	tgag	12258
12259	gata	cgcc	cgac	gtct	acag	ttgg	acta	9 9 9a	agac	ctct	aaca	gccc	cc								12308

GTATa at cactecc tett gagga acta et get et teac geagaa ag eget tag ceat gg eg tragt at gagt get geget et geget et geget et geget et geget gege

GTaatcactccctgtgaggaactactgtcttcacgcagaaagcgtctagccatggcgttagtatgagtgtcgtgcagcctccaggacccccctcccgggagagccatagtggtctgcggaaccggtgagtacaccggaattgccaggacgggtcctttcttggataaacccgctcaatgcctggagattggggtgccccgcaagactgctagccgagtagtgttgggtcgcgaaaggccttgtggtactgcctgagaggtgcttcgtagaccgtgcaccATG

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GTATTGCAGTTT gecage cect gat ggggg gacactecac cat gaat cactecc cet gt gaggaactact gtcttcac gcagaaagegtetagecatggegttagtatgagtgtegtgeagectecaggacceccctccegggagagecatagtggtetgeggaac cggtgagtacaccggaattgccaggacgaccgggtcctticttggataaacccgctcaatgcctggagatttgggcgtgccccgcaa gactgctagccgagtagtgttgggtcgcgaaaggccttgtggtactgcctgatagggtgcttgcgagtgccccgggaggtctcgtagaccgtgcaccATGGAGTTGATCACAAATGAACTTTTATACAAAACATACAAAAAC CČĞTCGGGGTGGAGGAACCTGTTTATGATCAGGCAGGTGATCCCTTATTTGGT GAAAGGGGAGCAGTCCACCCTCAATCGACGCTAAAGCTCCCACACAAGAGAG GGGAACGCGATGTTCCAACCAACTTGGCATCCTTACCAAAAAGAGGTGACTGC <u>AGGTCGGGTAATAGCAGAGGACCTGTGAGCGGGATCTACCTGAAGCCAGGGC</u> CACTATTTTACCAGGACTATAAAGGTCCCGTCTATCACAGGGCCCCGCTGGAGC TCTTTGAGGAGGGATCCATGTGTGAAACGACTAAACGGATAGGGAGAGTAACT ATAAAAGTGCCACGAGAAGTTACCAAAGGGTGTTCAGGTGGGTCCATAATAG GCTTGACTGCCTCTATGGGTCACAACTTGCTCAGACACGAAAGAAGAGGGAG CAACAAAAAGAAAACACAGAAACCCGACAGACTAGAAAGGGGGAAAATGAA AATAGTGCCCAAAGAATCTGAAAAAGACAGCAAAACTAAACCTCCGGATGCTA CAATAGTGGTGGAAGGAGTCAAATACCAGGTGAGGAAGAAGGGAAAAAACCAA CACGCAAGAACTGGAAAAAGCATTGTTGGCGTGGGCAATAATAGCTATAGTT TTGTTTCAAGTTACAATGGGAGAAAACATAACACAGTGGAACCTACAAGATAAT GGGACGGAAGGGATACAACGGGCAATGTTCCAAAGGGGTGTGAATAGAAGTT TACATGGAATCTGGCCAGAGAAAATCTGTACTGGTGTCCCTTCCCATCTAGCCA CCGATATAGAACTAAAAACAATTCATGGTATGATGGATGCAAGTGAGAAGACC CAACTGGTACAATATTGAACCCTGGATTCTAGTCATGAATAGAACCCAAGCCAA TCTCACTGAGGGACAACCACCAAGGGAGTGCGCAGTCACTTGTAGGTATGATA GGGCTAGTGACTTAAACGTGGTAACACAAGCTAGAGATAGCCCCACACCCTTA ACAGGTTGCAAGAAAGGAAAGAACTTCTCCTTTGCAGGCATATTGATGCGGGG CCCCTGCAACTTTGAAATAGCTGCAAGTGATGTATTATTCAAAGAACATGAACG CATTAGTATGTTCCAGGATACTACTCTTTACCTTGTTGACGGGTTGACCAACTCC TTAGAAGGTGCCAGACAAGGAACCGCTAAACTGACAACCTGGTTAGGCAAGCA GCTCGGGATACTAGGAAAAAGTTGGAAAACAAGAGTAAGACGTGGTTTGGAG CATACGCTGCTTCCCCTTACTGTGATGTCGATCGCAAAATTGGCTACATATGGT ATACAAAAATTGCACCCTGCCTGCTTACCCAAGAACACAAAAATTGTCGGCC CTGGGAAATTTGACACCAATGCAGAGGACGGCAAGATATTACATGAGATGGGG GGTCACTTGTCGGAGGTACTACTTTCTTTAGTGGTGCTGTCCGACTTCGCA CCGGAAACAGCTAGTGTAATGTACCTAATCCTACATTTTTCCATCCCACAAAGTC ACGTTGATGTAATGGATTGTGATAAGACCCAGTTGAACCTCACAGTGGAGCTG TATAAGACCAAATTGGTGGCCTTATGAGACAACTGTAGTGTTGGCATTTGAAGA GGTGAGCCAGGTGGTGAAGTTAGTGTTGAGGGCACTCAGAGATTTAACACGCA TTTGGAACGCTGCAACAACTACTGCTTTTTTAGTATGCCTTGTTAAGATAGTCAG GGGCCAGATGGTACAGGGCATTCTGTGGCTACTATTGATAACAGGGGTACAAG GGCACTTGGATTGCAAACCTGAATTCTCGTATGCCATAGCAAAGGACGAAAGA TGGAATGAAGCTGGAAGACACAATGGTCATTGCTTGGTGCGAAGATGGGAAGT TAATGTACCTCCAAAGATGCACGAGAGAAACCAGGTATCTCGCAATCTTGCATA CAAGAGCCTTGCCGACCAGTGTGGTATTCAAAAAACTCTTTGATGGGCGAAAG

CAAGAGGATGTAGTCGAAATGAACGACAACTTTGAATTTGGACTCTGCCCATGT GATGCCAAACCCATAGTAAGAGGGAAGTTCAATACAACGCTGCTGAACGGACC GGCCTTCCAGATGGTATGCCCCATAGGATGGACAGGGACTGTAAGCTGTACGT CATTCAATATGGACACCTTAGCCACAACTGTGGTACGGACATATAGAAGGTCTA AACCATTCCCTCATAGGCAAGGCTGTATCACCCAAAAGAATCTGGGGGAGGAT CTCCATAACTGCATCCTTGGAGGAAATTGGACTTGTGTGCCTGGAGACCAACTA CTATACAAAGGGGGCTCTATTGAATCTTGCAAGTGGTGTGGCTATCAATTTAAA GAGAGTGAGGGACTACCACACTACCCCATTGGCAAGTGTAAATTGGAGAACGA GACTGGTTACAGGCTAGTAGACAGTACCTCTTGCAATAGAGAAGGTGTGGCCA TAGTACCACAAGGGACATTAAAGTGCAAGATAGGAAAAACAACTGTACAGGTC ATAGCTATGGATACCAAACTCGGACCTATGCCTTGCAGACCATATGAAATCATA TCAAGTGAGGGCCTGTAGAAAAGACAGCGTGTACTTTCAACTACACTAAGAC ATTAAAAAATAAGTATTTTGAGCCCAGAGACAGCTACTTTCAGCAATACATGCT ATTACTTCGCTGAGTCCATATTAGTGGTGGTAGTAGCCCTCTTGGGTGGCAGAT ATGTACTTTGGTTACTGGTTACATACATGGTCTTATCAGAACAGAAGGCCTTAG GGATTCAGTATGGATCAGGGGAAGTGGTGATGATGGGCAACTTGCTAACCCAT AACAATATTGAAGTGGTGACATACTTCTTGCTGCTGTACCTACTGCTGAGGGAG GAGAGCGTAAAGAAGTGGGTCTTACTCTTATACCACATCTTAGTGGTACACCCA ATCAAATCTGTAATTGTGATCCTACTGATGATTGGGGATGTGGTAAAGGCCGAT TCAGGGGGCCAAGAGTACTTGGGGAAAATAGACCTCTGTTTTACAACAGTAGT ACTAATCGTCATAGGTTTAATCATAGCCAGGCGTGACCCAACTATAGTGCCACT GGTAACAATAATGGCAGCACTGAGGGTCACTGAACTGACCCACCAGCCTGGAG TTGACATCGCTGTGGCGGTCATGACTATAACCCTACTGATGGTTAGCTATGTGA CAGATTATTTTAGATATAAAAAATGGTTACAGTGCATTCTCAGCCTGGTATCTGC GGTGTTCTTGATAAGAAGCCTAATATACCTAGGTAGAATCGAGATGCCAGAGG TAACTATCCCAAACTGGAGACCACTAACTTTAATACTATTATATTTGATCTCAAC AACAATTGTAACGAGGTGGAAGGTTGACGTGGCTGGCCTATTGTTGCAATGTG TGCCTATCTTATTGCTGGTCACAACCTTGTGGGCCGACTTCTTAACCCTAATACT GATCCTGCCTACCTATGAATTGGTTAAATTATACTATCTGAAAACTGTTAGGACT GATATAGAAAGAAGTTGGCTAGGGGGGATAGACTATACAAGAGTTGACTCCAT CTACGACGTTGATGAGAGTGGAGAGGGCGTATATCTTTTTCCATCAAGGCAGA AAGCACAGGGGAATTTTTCTATACTCTTGCCCCTTATCAAAGCAACACTGATAA GTTGCGTCAGCAGTAAATGGCAGCTAATATACATGAGTTACTTAACTTTGGACT TTATGTACTACATGCACAGGAAAGTTATAGAAGAGATCTCAGGAGGTACCAACA TAATATCCAGGTTAGTGGCAGCACTCATAGAGCTGAACTGGTCCATGGAAGAA GAGGAGAGCAAAGGCTTAAAGAAGTTTTATCTATTGTCTGGAAGGTTGAGAAA CCTAATAATAAAACATAAGGTAAGGAATGAGACCGTGGCTTCTTGGTACGGGG AGGAGGAAGTCTACGGTATGCCAAAGATCATGACTATAATCAAGGCCAGTACA CTGAGTAAGAGCAGGCACTGCATAATATGCACTGTATGTGAGGGCCGAGAGTG GAAAGGTGGCACCTGCCCAAAATGTGGACGCCATGGGAAGCCGATAACGTGT GGGATGTCGCTAGCAGATTTTGAAGAAAGACACTATAAAAGAATCTTTATAAGG GAAGGCAACTTTGAGGGTATGTGCAGCCGATGCCAGGGAAAGCATAGGAGGT TTGAAATGGACCGGGAACCTAAGAGTGCCAGATACTGTGCTGAGTGTAATAGG CTGCATCCTGCTGAGGAAGGTGACTTTTGGGCAGAGTCGAGCATGTTGGGCCT CAAAATCACCTACTTTGCGCTGATGGATGGAAAGGTGTATGATATCACAGAGTG GGCTGGATGCCAGCGTGTGGGAATCTCCCCAGATACCCACAGAGTCCCTTGTC ACATCTCATTTGGTTCACGGATGCCTTTCAGGCAGGAATACAATGGCTTTGTAC AATATACCGCTAGGGGGCAACTATTTCTGAGAAACTTGCCCGTACTGGCAACTA AAGTAAAAATGCTCATGG1AGGCAACCTTGGAGAAGAAATTGGTAATCTGGAA CATCTTGGGTGGATCCTAAGGGGGCCTGCCGTGTGTAAGAAGATCACAGAGCA CGAAAAATGCCACATTAATATACTGGATAAACTAACCGCATTTTTCGGGATCAT GCCAAGGGGACTACACCCAGAGCCCCGGTGAGGTTCCCTACGAGCTTACTAA

AAGTTCAGTCGACCATGTAACCGCCGGAAAAGATCTACTGGTCTGTGACAGCA TGGGACGAACTAGAGTGGTTTGCCAAAGCAACAACAGGTTGACCGATGAGACA GAGTATGGCGTCAAGACTGACTCAGGGTGCCCAGACGGTGCCAGATGTTATGT GTTAAATCCAGAGGCCGTTAACATATCAGGATCCAAAGGGGCAGTCGTTCACC TCCAAAAGACAGGTGGAGAATTCACGTGTGTCACCGCATCAGGCACACCGGCT TTCTTCGACCTAAAAAACTTGAAAGGATGGTCAGGCTTGCCTATATTTGAAGCC TCCAGCGGGAGGGTTGGTTGGCAGAGTCAAAGTAGGGAAGAATGAAGAGTCTA AACCTACAAAAATAATGAGTGGAATCCAGACCGTCTCAAAAAACAGAGCAGAC CTGACCGAGATGGTCAAGAAGATAACCAGCATGAACAGGGGAGACTTCAAGCA GATTACTTTGGCAACAGGGCAGGCAAAACCACAGAACTCCCAAAAGCAGTTA TAGAGGAGATAGGAAGACACAAGAGAGTATTAGTTCTTATACCATTAAGGGCA GCGGCAGAGTCAGCCAGTATATGAGATTGAAACACCCAAGCATCTCTTTT AACCTAAGGATAGGGGACATGAAAGAGGGGGACATGGCAACCGGGATAACCT ATGCATCATACGGGTACTTCTGCCAAATGCCTCAACCAAAGCTCAGAGCTGCTA TGGTAGAATACTCATACATATTCTTAGATGAATACCATTGTGCCACTCCTGAACA ACTGGCAATTATCGGGAAGATCCACAGATTTTCAGAGAGTATAAGGGTTGTCG CCATGACTGCCACGCCAGCAGGGTCGGTGACCACAACAGGTCAAAAGCACCCA ATAGAGGAATTCATAGCCCCCGAGGTAATGAAAGGGGAGGATCTTGGTAGTCA GTTCCTTGATATAGCAGGGTTAAAAATACCAGTGGATGAGATGAAAGGCAATAT GTTGGTTTTTGTACCAACGAGAAACATGGCAGTAGAGGTAGCAAAGAAGCTAA AAGCTAAGGGCTATAACTCTGGATACTATTACAGTGGAGAGGATCCAGCCAATCTGAGAGTTGTGACATCACAATCCCCCTATGTAATCGTGGCTACAAATGCTATT GAATCAGGAGTGACACTACCAGATTTGGACACGGTTATAGACACGGGGTTGAA ATGTGAAAAGAGGGTGAGGGTATCATCAAAGATACCCTTCATCGTAACAGGCC TTAAGAGGATGGCCGTGACTGTGGGTGAGCAGGCGCAGCGTAGGGGCAGAGT AGGTAGAGTGAAACCCGGGAGGTATTATAGGAGCCAGGAAACAGCAACAGGG TCAAAGGACTACCACTATGACCTCTTGCAGGCACAAAGATACGGGATTGAGGA TGGAATCAACGTGACGAAATCCTTTAGGGAGATGAATTACGATTGGAGCCTATA CGAGGAGGACAGCCTACTAATAACCCAGCTGGAAATACTAAATAATCTACTCAT CTCAGAAGACTTGCCAGCCGCTGTTAAGAACATAATGGCCAGGACTGATCACC CAGAGCCAATCCAACTTGCATACAACAGCTATGAAGTCCAGGTCCCGGTCCTGT TCCCAAAAATAAGGAATGGAGAAGTCACAGACACCTACGAAAATTACTCGTTTC TAAATGCCAGAAAGTTAGGGGAGGATGTGCCCGTGTATATCTACGCTACTGAA GATGAGGATCTGGCAGTTGACCTCTTAGGGCTAGACTGGCCTGATCCTGGGAA CCAGCAGGTAGTGGAGACTGGTAAAGCACTGAAGCAAGTGACCGGGTTGTCCT CGGCTGAAAATGCCCTACTAGTGGCTTATTTGGGTATGTGGGTTACCAGGCTC TCTCAAAGAGGCATGTCCCAATGATAACAGACATATATACCATCGAGGACCAGA GACTAGAAGACACCACCCACCTCCAGTATGCACCCAACGCCATAAAAACCGAT GGGACAGAGACTGAAAGAACTGGCGTCGGGTGACGTGGAAAAAATCA TGGGAGCCATTTCAGATTATGCAGCTGGGGGACTGGAGTTTGTTAAATCCCAA GCAGAAAAGATAAAAACAGCTCCTTTGTTTAAAGAAAACGCAGAAGCCGCAAA AGGGTATGTCCAAAAATTCATTGACTCATTAATTGAAAATAAAGAAGAAATAAT CAGATATGGTTTGTGGGGAACACACACAGCACTATACAAAAGCATAGCTGCAA GACTGGGGCATGAAACAGCGTTTGCCACACTAGTGTTAAAGTGGCTAGCTTTT GGAGGGGAATCAGTGTCAGACCACGTCAAGCAGGCGGCAGTTGATTTAGTGG TCTATTATGTGATGAATAAGCCTTCCTTCCCAGGTGACTCCGAGACACAGCAAG AAGGGAGGCGATTCGTCGCAAGCCTGTTCATCTCCGCACTGGCAACCTACACA TACAAAACTTGGAATTACCACAATCTCTCTAAAGTGGTGGAACCAGCCCTGGCT TACCTCCCCTATGCTACCAGCGCATTAAAAATGTTCACCCCAACGCGGCTGGAG AGCGTGGTGATACTGAGCACCACGATATATAAAACATACCTCTCTATAAGGAAG GGGAAGAGTGATGGATTGCTGGGTACGGGGATAAGTGCAGCCATGGAAATCC TGTCACAAAACCCAGTATCGGTAGGTATATCTGTGATGTTGGGGGTAGGGGCA ATCGCTGCGCACAACGCTATTGAGTCCAGTGAACAGAAAAGGACCCTACTTAT GAAGGTGTTTGTAAAGAACTTCTTGGATCAGGCTGCAACAGATGAGCTGGTAA

AAGAAAACCCAGAAAAATTATAATGGCCTTATTTGAAGCAGTCCAGACAATTG GTAACCCCTGAGACTAATATACCACCTGTATGGGGTTTACTACAAAGGTTGGG AGGCCAAGGAACTATCTGAGAGGACAGCAGGCAGAAACTTATTCACATTGATA ATGTTTGAAGCCTTCGAGTTATTAGGGATGGACTCACAAGGGAAAATAAGGAA CCTGTCCGGAAATTACATTTTGGATTTGATATACGGCCTACACAAGCAAATCAA CAGAGGCTGAAGAAAATGGTACTGGGGTGGGCCCCTGCACCCTTTAGTTGTG ACTGGACCCCTAGTGACGAGAGGATCAGATTGCCAACAGACAACTATTTGAGG GTAGAAACCAGGTGCCCATGTGGCTATGAGATGAAAGCTTTCAAAAATGTAGG TGGCAAACTTACCAAAGTGGAGGAGAGCGGGCCTTTCCTATGTAGAAACAGAC CTGGTAGGGGACCAGTCAACTACAGAGTCACCAAGTATTACGATGACAACCTC AGAGAGATAAAACCAGTAGCAAAGTTGGAAGGACAGGTAGAGCACTACTACAA AGGGGTCACAGCAAAAATTGACTACAGTAAAGGAAAAATGCTCTTGGCCACTG ACAAGTGGGAGGTGGAACATGGTGTCATAACCAGGTTAGCTAAGAGATATACT GGGGTCGGGTTCAATGGTGCATACTTAGGTGACGAGCCCAATCACCGTGCTCT AGTGGAGAGGGACTGTGCAACTATAACCAAAAACACAGTACAGTTTCTAAAAAT GAAGAAGGGTGTGCGTTCACCTATGACCTGACCATCTCCAATCTGACCAGGC TCATCGAACTAGTACACAGGAACAATCTTGAAGAGAAGGAAATACCCACCGCT ACGGTCACCACATGGCTAGCTTACACCTTCGTGAATGAAGACGTAGGGACTAT AAAACCAGTACTAGGAGAGAGAGTAATCCCCGACCCTGTAGTTGATATCAATTT ACAACCAGAGGTGCAAGTGGACACGTCAGAGGTTGGGATCACAATAATTGGAA GGGAAACCCTGATGACAACGGGAGTGACACCTGTCTTGGAAAAAGTAGAGCCT GACGCCAGCGACAACCAAAACTCGGTGAAGATCGGGTTGGATGAGGGTAATTA CCCAGGGCCTGGAATACAGACACATACACTAACAGAAGAAATACACAACAGGG ATGCGAGGCCCTTCATCATGATCCTGGGCTCAAGGAATTCCATATCAAATAGGG CAAAGACTGCTAGAAATATAAATCTGTACACAGGAAATGACCCCAGGGAAATA CGAGACTTGATGGCTGCAGGGCGCATGTTAGTAGTAGCACTGAGGGATGTCGA CCTGAGCTGTCTGAAATGGTCGATTTCAAGGGGACTTTTTTAGATAGGGAGG CCCTGGAGGCTCTAAGTCTCGGGCAACCTAAACCGAAGCAGGTTACCAAGGAA GCTGTTAGGAATTTGATAGAACAGAAAAAGATGTGGAGATCCCTAACTGGTTT GCATCAGATGACCCAGTATTTCTGGAAGTGGCCTTAAAAAAATGATAAGTACTAC TTAGTAGGAGATGTTGGAGAGCTAAAAGATCAAGCTAAAGCACTTGGGGCCAC GGATCAGACAAGAATTATAAAGGAGGTAGGCTCAAGGACGTATGCCATGAAGC TATCTAGCTGGTTCCTCAAGGCATCAAACAAACAGATGAGTTTAACTCCACTGT TTGAGGAATTGTTGCTACGGTGCCCACCTGCAACTAAGAGCAATAAGGGGCAC ATGGCATCAGCTTACCAATTGGCACAGGGTAACTGGGAGCCCCTCGGTTGCGG GGTGCACCTAGGTACAATACCAGCCAGAAGGGTGAAGATACACCCATATGAAG CTTACCTGAAGTTGAAAGATTTCATAGAAGAAGAAGAAGAAACCTAGGGTT CAAGGAAACCTCAACACCAAGAAAATGCTCAACCCGGGGAAACTATCTGAACA **GTTGGACAGGGAGGGGCGCAAGAGGAACATCTACAACCACCAGATTGGTACT** ATAATGTCAAGTGCAGGCATAAGGCTGGAGAAATTGCCAATAGTGAGGGCCCA **AACCGACACAAACCTTTCATGAGGCAATAAGAGATAAGATAGACAAGAGTG** AAAACCGGCAAAATCCAGAATTGCACAACAAATTGTTGGAGATTTTCCACACGA TAGCCCAACCCACCTGAAACACACCTACGGTGAGGTGACGTGGGAGCAACTT GAGGCGGGATAAATAGAAAGGGGGCAGCAGGCTTCCTGGAGAAGAAGAACA TCGGAGAAGTATTGGATTCAGAAAAGCACCTGGTAGAACAATTGGTCAGGGAT CTGAAGGCCGGGAGAAAGATAAAATATTATGAAACTGCAATACCAAAAAATGA GAAGAGAGATGTCAGTGATGACTGGCAGGCAGGGGACCTGGTGGTTGAGAAG AGGCCAAGAGTTATCCAATACCCTGAAGCCAAGACAAGGCTAGCCATCACTAA GGTCATGTATAACTGGGTGAAACAGCAGCCCGTTGTGATTCCAGGATATGAAG GAAAGACCCCCTTGTTCAACATCTTTGATAAAGTGAGAAAGGAATGGGACTCGT TCAATGAGCCAGTGGCCGTAAGTTTTGACACCAAAGCCTGGGACACTCAAGTG ACTAGTAAGGATCTGCAACTTATTGGAGAAATCCAGAAATATTACTATAAGAAG GAGTGGCACAAGTTCATTGACACCATCACCGACCACATGACAGAAGTACCAGT

TATAACAGCAGATGGTGAAGTATATATAAGAAATGGGCAGAGAGGGAGCGGC CAGCCAGACACAGTGCTGGCAACAGCATGTTAAATGTCCTGACAATGATGTA CGGCTTCTGCGAAAGCACAGGGGTACCGTACAAGAGTTTCAACAGGGTGGCAA GGATCCACGTCTGTGGGGATGATGGCTTCTTAATAACTGAAAAAGGGTTAGGG CTGAAATTTGCTAACAAAGGGATGCAGATTCTTCATGAAGCAGGCAAACCTCAG AAGATAACGGAAGGGGAAAAGATGAAAGTTGCCTATAGATTTGAGGATATAGA **GTTCTGTTCTCATACCCCAGTCCCTGTTAGGTGGTCCGACAACACCAGTAGTCA** CATGGCCGGGAGAGACACCGCTGTGATACTATCAAAGATGGCAACAAGATTGG ATTCAAGTGGAGAGAGGGGTACCACAGCATATGAAAAAGCGGTAGCCTTCAGT TTCTTGCTGATGTATTCCTGGAACCCGCTTGTTAGGAGGATTTGCCTGTTGGTC CTTTCGCAACAGCCAGAGACAGACCCATCAAAACATGCCACTTATTATTACAAA GGTGATCCAATAGGGGCCTATAAAGATGTAATAGGTCGGAATCTAAGTGAACT GAAGAGAACAGGCTITGAGAAATTGGCAAATCTAAACCTAAGCCTGTCCACGTT GGGGATCTGGACTAAGCACACAAGCAAAAGAATAATTCAGGACTGTGTTGCCA TTGGGAAAGAAGAGGCCAACTGGCTAGTTAACGCCGACAGGCTGATATCCAGC AAAACTGGCCACTTATACATACCTGATAAAGGCTTTACATTACAAGGAAAGCAT TATGAGCAACTGCAGCTAAGAACAGAGACAAACCCGGTCATGGGGGTTGGGA CTGAGAGATACAAGTTAGGTCCCATAGTCAATCTGCTGCTGAGAAGGTTGAAA **ATTCTGCTCATGACGGCCGTCGGCGTCAGCAGCTGAaggttggggtaaacactccggcctcttag** gecameetgummuummuummuummuummuummuumeetmuummuumittettieettettiiteettettiite cttcettctttaatggtggctccatcttagccctagtcacggctagctgtgaaaggtccgtgagccgcatgactgcagagagtgctgatactggcctctctgcagatcatgtCCCCGGCCGTCGGCGTCAGCTGAgacaaaatgtatatattgtaaataaattaatc catgtacatagtgtatataaatatagttgggaccgtccacctcaagaagacgacacgcccaacacgcacagctaaacagtagtcaagatt atctacctcaagataacactacatttaatgcacacagcactttagctgtatgaggatacgcccgacgtctatagttggactagggaagacct ctaacagccccc

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		10	20				CCGGCCTCT	MAGGCCA
3H3Bfrag	\ \ \ mm~~~~			30	1 40	50	60	70
	AATICIG	CICAIGA	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	CCCTCACCAC	TGAAGGTYG	CATABACAC		• •
1.1.4 seq								
1.2.3 seq	AATTCTG	CTCATGAC	CGCCCGTYC	CONCION		COLAMALAL!	CCGGCCICT	MAGGCCA 70
6.2.2 seq	AATTOTO	TATEA		GCGTCAGCAG	TOMOGING	AGTAAACAC!	CCCCCCTCTT	MAGGCCA 70
6,1.4 seg								
Train seq	WIICIG	CICHIGH	-GCCCG1CG	CCCTCACCAC	TGAAGGTTGC	GGTAAACACT	CCGGCCTCTT	AGGCCA 70
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	TTTCCTG	THEFT	MAMMAMA	باململململململ				
21275		. 80	90	100	110	120	130	140
3H3Bfrag	TITICCIG	TTTTTTT	MALABATA		Alalalalalalalalalalalalalalalalalalala	<u>-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1</u>		140
1.1.4 seq	TTTCCTG	TITTITI	MANANANI	TTTTTTTTT	atatatata() [1111111111	TITICITIT	
1;2.3 seq	TTTCCTG	PERMIT	TITTTTT	Tatatalaisisis				109
6,2.2 seq	Tabab Ashab.	<u> </u>	TITITIT		~			102
6.1.4 seq	TTTCCTG		11111111	11111				99
vier seq	TITCE IG.	TTTTT		:				04
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		250			TATCHARAIA	CCLLICIALI	CCTTCCTTCT	TTAATG .
211275		150	160	170	180	190	200	210
3H3Bfrag	TITTTTCC	CITITITI	TITITITI	TTTTCTTTCC	MAINTERPRETATION	~	~~~~	210
1 1.4 seq						CTITCITIN	CTICCTICT	PTAATG 210
1.12.3 seq					TACTTTTT-	CLITCITTI	CTICCTICT.	TTANDS 149
6.12.2 seg				CTTICC	TICTITITI-	CTITCTTFR	CTICCTICT	TTAATG 142
6.1.4 seq				CITICC	PECTETER	CLERCLANA	ملم فلملم في الملم في	PRANTO 140
				CTTTCC	TTCTTTTTTT	Medalah Malalad	Andrew Mark	Market 100
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	GTGGCTCC	ATCTTAC	CTACTCA					
	GTGGCTCC	ATCTTAC	CCCTAGTCA	CGGCTAGCTG				SAGTGC
31725	•	220	230	CGGCTAGCTG	TGANAGGTCCC	TCAGCCGCA	GACTGCAGAC	
3H3Bfrag	GTGGCTCC	220 ATCITAGO	230 CCTAGTCA	CGCTAGCTG 240 CGCTACCTG	TGAAAGGTCCC	ZGO	CACTGCAGAC 270	280
1.1.4 seq	GIGGCICC	220 ATCITAGO ATCITAGO	UES CCTAGTCA CCTAGTCA	CGCTACCTG 240 CGCTACCTG	TGAAAGGTCCC 250 TGAAAGGTCCC	FIGAGCOCA 260 FIGAGCOCAI	IGACTICCAGAC 270 IGACTICCAGAC	280 SAGTGC 280
1.1.4 seq 1.2.3 seq	GIGGCICC	220 ATCITAGO ATCITAGO	UES CCTAGTCA CCTAGTCA	CGCTACCTG 240 CGCTACCTG	TGAAAGGTCCC 250 TGAAAGGTCCC	FIGAGCOCA 260 FIGAGCOCAI	IGACTICCAGAC 270 IGACTICCAGAC	280 SAGTGC 280
1.1.4 seq 1.2.3 seq	GIGGCICC GIGGCICC	220 ATCTTAGO ATCTTAGO	UES COTAGICA COTAGICA COTAGICA	CGGCTAGCTG 240 CGGCTAGCTG CGGCTAGCTG	TGAAAGGTCCC 250 TGAAAGGTCCC	Z60 STGAGCCGCAI STGAGCCGCAI	GACTGCAGAC 270 GACTGCAGAC GACTGCAGAC	280 PAGTGC 280 PAGTGC 219
1.11.4 seq 1.2.3 seq 6.2.2 seq	GIGGCICC GIGGCICC GIGGCICC	220 ATCTTAGO ATCTTAGO ATCTTAGO	DES COTAGICA COTAGICA COTAGICA COTAGICA	CGGCTAGCTG 240 CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG	TGAAAGGTCCC 250 TGAAAGGTCCC TGAAAGGTCCC	260 FTGAGCCGCAI FTGAGCCGCAI FTGAGCCGCAI	GACTGCAGAC 270 GACTGCAGAC GACTGCAGAC	280 SAGTOC 280 SAGTOC 219 SAGTOC 212
1.1.4 seq 1.2.3 seq	GIGGCICC GIGGCICC GIGGCICC	220 ATCTTAGO ATCTTAGO ATCTTAGO	DES COTAGICA COTAGICA COTAGICA COTAGICA	CGGCTAGCTG 240 CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG	TGAAAGGTCCC 250 TGAAAGGTCCC TGAAAGGTCCC	260 FTGAGCCGCAI FTGAGCCGCAI FTGAGCCGCAI	GACTGCAGAC 270 GACTGCAGAC GACTGCAGAC	280 SAGTOC 280 SAGTOC 219 SAGTOC 212
1.11.4 seq 1.2.3 seq 6.2.2 seq	GIGGCICC GIGGCICC GIGGCICC	220 ATCTTAGO ATCTTAGO ATCTTAGO	DES COTAGICA COTAGICA COTAGICA COTAGICA	CGGCTAGCTG 240 CGGCTAGCTG CGGCTAGCTG	TGAAAGGTCCC 250 TGAAAGGTCCC TGAAAGGTCCC	260 FTGAGCCGCAI FTGAGCCGCAI FTGAGCCGCAI	GACTGCAGAC 270 GACTGCAGAC GACTGCAGAC	280 SAGTOC 280 SAGTOC 219 SAGTOC 212
1.11.4 seq 1.2.3 seq 6.2.2 seq	GTGGCTCC GTGGCTCC GTGGCTCC	ATCTTAGG ATCTTAGG ATCTTAGG ATCTTAGG ATCTTAGG	230 CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA	CGGCTAGCTG 240 CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG	TCAAAGGTCCC 250 TCAAAGGTCCCC TCAAAGGTCCCC TCAAAGGTCCCC TCAAAGGTCCCC	260 260 STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN	TEACTIGCAGAC 270 TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC	280 SAGTOC 280 SAGTOC 219 SAGTOC 212 SAGTOC 210 SAGTOC 195
1.11.4 seq 1.2.3 seq 6.2.2 seq	GTGGCTCC GTGGCTCC GTGGCTCC	ATCTTAGG ATCTTAGG ATCTTAGG ATCTTAGG ATCTTAGG	230 CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA	CGGCTAGCTG 240 CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG	TCAAAGGTCCC 250 TCAAAGGTCCCC TCAAAGGTCCCC TCAAAGGTCCCC TCAAAGGTCCCC	260 260 STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN	TEACTIGCAGAC 270 TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC	280 SAGTOC 280 SAGTOC 219 SAGTOC 212 SAGTOC 210 SAGTOC 195
1.11.4 seq 1.2.3 seq 6.2.2 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC	ATCTTAGE ATCTTAGE ATCTTAGE ATCTTAGE ATCTTAGE ATCTTAGE	230 CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA	CGGCTAGCTG 240 CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG	TGAAAGGTCCC 250 TGAAAGGTCCC TGAAAGGTCCC TGAAAGGTCCC TGAAAGGTCCC	FIGAGCOGCAY 260 FIGAGCOGCAY FIGAGCOGCAY FIGAGCOGCAY FIGAGCCOCAY FIGAGCCOCAY	TEACTIGCAGAC 270 TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC	280 SAGTOC 280 SAGTOC 219 SAGTOC 212 SAGTOC 210 SAGTOC 195
1.11.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC	ATCTTAGE ATC	230 CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA TGCAGATCA	CGGCTAGCTG 240 CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG	TCAAAGGTCCC 250 TCAAAGGTCCC TCAAAGGTCCC TCAAAGGTCCC TCAAAGGTCCC	260 27GAGCCGCAN 27GAGCCGCAN 27GAGCCGCAN 27GAGCCGCAN 27GAGCCGCAN 27GAGCCGCAN 27GAGCCGCAN 27GAGCCGCAN	TEACTIGCAGAC 270 TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC	280 SAGTOC 280 SAGTOC 219 SAGTOC 212 SAGTOC 210 SAGTOC 195
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 3H3Bfrag	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC	ATCTTAGE ATC	230 CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA TGCAGATCA TGCAGATCA	CGGCTAGCTG 240 CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG	TCAAAGGTCCC 250 TCAAAGGTCCCC TCAAAGGTCCCCT TCAAAAGGTCCCCT TCAAAAGGTCCCT TCAAAAGGTCCCCT TCAAAAGGTCCCCT TCAAAAGGTCCCCT TCAAAAGGTCCCCT TCAAAAGGTCCCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCCT TCAAAAGGTCCT TCAAAAGGTCCT TCAAAAGGTCCT TCAAAAGGTCCT TCAAAAGGTCCT TCAAAAGGTCCT TCAAAAGGTCCT TCAAAAAGGTCCT TCAAAAGGTCCT TCAAAAGGTCCT TCAAAAGGTCCT TCAAAAAGGTCCT TCAAAAAGGTCCT TCAAAAAGGTCCT TCAAAAAGGTCCT TCAAAAGGTCCT TCAAAAAGGTCCT TCAAAAAAGGTCCT TCAAAAAAGGTCCT TCAAAAAAGGTCCT TCAAAAAAGGTCCT TCAAAAAAGGTCCT TCAAAAAAGGTCCT TCAAAAAAAGGTCCT TCAAAAAAGGTCCT TCAAAAAAAGGTCCT TCAAAAAAAGGTCCT TCAAAAAAAAAA	260 STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN	TEACTIGCAGAC 270 TEACTIGCAGAC TEACTIGCACAC TEACTIGCAGAC TEACTIC TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC TEACTIGCAGAC TEACTIGCACAC TEACTICCAGAC TEACTICCAGAC TEACTICCAGAC TEACTICCAGAC TEACTICCAGAC TEACTICCAGAC TEACTICCAGAC TEACTICCAGAC TEACTICCACAC TEACTICCACAC TEACTICCACAC TEACTICCACAC TEACTICCACAC TEACTICCACAC TEACT	280 SAGTOC 280 SAGTOC 219 SAGTOC 212 SAGTOC 210 SAGTOC 195 SAGTOC 350
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 3H3Bfrag 1.1.4 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC TGATACTC	ATCTTAGE ATCTTTAGE ATCTTTAG	230 230 2CCTAGTCA 2CCTAGTCA 2CCTAGTCA 2CCTAGTCA 2CCTAGTCA 3CCTAGTCA 300 300 300 300 300 300 300 300 300 30	CGCTAGCTG 240 CGCTAGCTG CGCCTAGCTG CGCTAGCTG CGCCTAGCTG CGCCTAGCTG CGCCTAGCTG CGCCTAGCTG CGCCTAG	TGAAAGGTCCC 250 TGAAAGGTCCCC	260 260 STGAGCCGCAN STGAGCAGCTGAG	TEACTICAGAE 270 TEACTICAGAE TEACTICAGAE TEACTICAGAE TEACTICAGAE TEACTICAGAE TEACTICAGAE TEACTICAGAE TEACTICAGAE TEACTICAGAE TEACATATOTA 340	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 PATAT 350 PATAT 350
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.1.4 seq 1.2.3 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC TGATACTC TGATACTC	ATCTTACE ATC	230 CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA TGCAGATCA TGCAGATCA TGCAGATCA	CGGCTAGCTG 240 CGGCTAGCTG C	TGAAAGGTCCC 250 TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCCCCCCCC	260 STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN AGCAGCTGAG	TEACTICACACA 270 TEACTICACACACACACACACACACACACACACACACACACA	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 PATAT 350 PATAT 350 PATAT 289
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC TGATACTC TGATACTC	ATCTTACE ATC	230 CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA TGCAGATCA TGCAGATCA TGCAGATCA	CGGCTAGCTG 240 CGGCTAGCTG C	TGAAAGGTCCC 250 TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCCCCCCCC	260 STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN STGAGCCGCAN AGCAGCTGAG	TEACTICACACA 270 TEACTICACACACACACACACACACACACACACACACACACA	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 PATAT 350 PATAT 350 PATAT 289
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC TGATACTC TGATACTC TGATACTC TGATACTC	ATCTTAGE ATC	230 CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA	CGGCTAGCTG 240 CGGCTAGCTG C	TGAAAGGTCCC 250 TGAAAGGTCCC TGAAAGGTCCCC TGAAAAGGTCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCCC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCTTC TGAAAAGGTCCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCCTTC TGAAAAGGTCCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAAGGTCCCTTC TGAAAAAAGGTCCCTTC TGAAAAAAGGTCCCTTC TGAAAAAAAGGTCCCTTC TGAAAAAAAGGTCCCTTC TGAAAAAAAAAA	260 STGAGCCGCAY TGAGCCGCAY STGAGCCGCAY STGAGCCGCAY STGAGCCGCAY AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG	TGACTGCAGAC 270 TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC ACAAAATGTA ACAAAATGTA ACAAAATGTA	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 TATAT 350 TATAT 289 TATAT 282
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.1.4 seq 1.2.3 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC TGATACTC TGATACTC TGATACTC TGATACTC	ATCTTAGE ATC	230 CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA	CGGCTAGCTG 240 CGGCTAGCTG C	TGAAAGGTCCC 250 TGAAAGGTCCC TGAAAGGTCCCC TGAAAAGGTCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCCC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCTTC TGAAAAGGTCCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCCTTC TGAAAAGGTCCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAAGGTCCCTTC TGAAAAAAGGTCCCTTC TGAAAAAAGGTCCCTTC TGAAAAAAAGGTCCCTTC TGAAAAAAAGGTCCCTTC TGAAAAAAAAAA	260 STGAGCCGCAY TGAGCCGCAY STGAGCCGCAY STGAGCCGCAY STGAGCCGCAY AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG	TCACTICCACAC 270 TCACTICCACAC TCACTICCACACAC TCACTICCACACACACACACACACACACACACACACACAC	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 TATAT 350 TATAT 289 TATAT 289
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC TGATACTC TGATACTC TGATACTC TGATACTC	ATCTTAGE ATC	230 CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA CCCAGATCA	CGGCTAGCTG 240 CGGCTAGCTG C	TGAAAGGTCCC 250 TGAAAGGTCCC TGAAAGGTCCCC TGAAAAGGTCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCC TGAAAAGGTCCCCCC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCTTC TGAAAAGGTCCCCCCTTC TGAAAAGGTCCCCCTTC TGAAAAGGTCCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCCTTC TGAAAAGGTCCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAGGTCCCTTC TGAAAAAAGGTCCCTTC TGAAAAAAGGTCCCTTC TGAAAAAAGGTCCCTTC TGAAAAAAAGGTCCCTTC TGAAAAAAAGGTCCCTTC TGAAAAAAAAAA	260 STGAGCCGCAY TGAGCCGCAY STGAGCCGCAY STGAGCCGCAY STGAGCCGCAY AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG	TCACTICCACAC 270 TCACTICCACAC TCACTICCACACAC TCACTICCACACACACACACACACACACACACACACACAC	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 TATAT 350 TATAT 289 TATAT 282
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGATACTC TGATACTC TGATACTC TGATACTC	220 ATCTTAGE	230 CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA TGCAGATCA TGCAGATCA TGCAGATCA TGCAGATCA TGCAGATCA	CGCTACTO 240 CGCTACTO C	TGAAAGGTCCC 250 TGAAAGGTCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGGAAAGGTCCCC TGGAAAGGTCCCC TGGAAAGGTCCCC TGGAAAGGTCCCC TGGAAAGGTCCCCTCCCC	260 FIGAGCOGCAN FIGAGCOGCAN FIGAGCCGCAN FIGAGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG	TCACTICCACAC 270 TCACTICCACAC TCACTICCACACAC TCACTICCACACACACACACACACACACACACACACACAC	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 TATAT 350 TATAT 289 TATAT 282
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGATACTC TGATACTC TGATACTC TGATACTC	220 ATCTTAGE	230 CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA TGCAGATCA TGCAGATCA TGCAGATCA TGCAGATCA TGCAGATCA	CGCTACTO 240 CGCTACTO C	TGAAAGGTCCC 250 TGAAAGGTCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGAAAGGTCCCC TGGAAAGGTCCCC TGGAAAGGTCCCC TGGAAAGGTCCCC TGGAAAGGTCCCC TGGAAAGGTCCCCTCCCC	260 FIGAGCOGCAN FIGAGCOGCAN FIGAGCCGCAN FIGAGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG	TCACTICCACAC 270 TCACTICCACAC TCACTICCACACAC TCACTICCACACACACACACACACACACACACACACACAC	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 TATAT 350 TATAT 289 TATAT 289
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGATACTG TGATACTG TGATACTG TGATACTG TGATACTG TGATACTG TGATACTG	ATTTAATC	CCTAGTCA CCTAGTCA CCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCTAGTCA CCCAGATCA CCCATGTACA CCCAGTTACA CCCAGATCA	CGGCTAGCTG 240 CGGCTAGCTG CGGCCCCGGC CGGCCCCGGC CGGCCCGGC CGGCCCGGC CGGCCCGGC CGGCCCGGC CGGCCCGGC CGGCCCGGCC CGGCCCGCC CGGCTAGCTG CGGCTAG	TGAAAGGTCCC 250 TGAAAGGTCCCC TGAAAGGTCCCCTCCCC	260 FIGAGCOGCAN FIGAGCOGCAN FIGAGCCGCAN FIGAGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG	TCACTICCACAC 270 TCACTICCACAC TCACTICCACACAC TCACTICCACACACACACACACACACACACACACACACAC	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 TATAT 350 TATAT 289 TATAT 282
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGATACTG TGATACTG TGATACTG TGATACTG TGATACTG TGATACTG	ATTTAATC	230 CCTAGTCA CCTAGTCA CCTAGTCA CCCTAGTCA CCCTAGTCA 300 CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA	CGGCTAGCTG 240 CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG CGGCTAGCTG TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC	TCAAAGGTCCC 250 TCAAAGGTCCC TCAAAAGGTCCC TCAAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCCC TCAAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAGGTCCC TCAAAAAGGTCCC TCAAAAAAAAGGTCCC TCAAAAAAAAAA	FIGAGCOGCAM 260 FIGAGCOGCAM FIGAGCOGCAM	TCACTICCACAC 270 TCACTICCACAC TCACTICCACACAC TCACTICCACACACACACACACACACACACACACACACAC	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 TATAT 350 TATAT 289 TATAT 282
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq 6.2.2 seq 6.1.4 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC TGATACTC	ATTTAATC	230 CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCAGATCA	CGCTAGCTG 240 CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC	TGAAAGGTCCC 250 TGAAAGGTCCC TGAAAGGTCCCC TCGTCGGCGTCCCCCTCCCCCTCCCCCTCCCCCTCCCCCTCCCCCTCCCC	260 STGAGCCGCAN STGAGCAGCTGAG AGCAGCTGAG	TCACTICCACAC 270 TCACTICCACAC TCACTICCACACAC TCACTICCACACACACACACACACACACACACACACACAC	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 195 PAGTOC 195 PAGTAT 350 PATAT 289 PATAT 289 PATAT 280 PATAT 280 PATAT 280 PATAT 280 PATAT 265
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.2.3 seq 6.1.4 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC TGATACTC TGTAAATAI	ATTTAATC ATTTAAC ATCTTAC ATTTAATC ATTTAATC ATTTAATC ATTTAATC ATTTAATC ATTTAATC ATTTAATC	230 CCTAGTCA CCAGATCA	CGCTAGCTG 240 CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG TGTCCCCCGG TGTCCCCCGG TGTCCCCGGG TGTCCCCCGGG TGTCCCCGGG TGTCCCCGGG TGTCCCCCGGG TGTCCCCCGGG TGTCCCCCGGG TGTCCCCCGGG TGTCCCCGGG TGTCCCCCGGG TGTCCCCCGG	TGAAAGGTCCC 250 TGAAAGGTCCC TGAAAAGGTCCC TGAAAAGGTCCCC TGAAAAGGTCCCC TGAAAAGGTCCCC TGAAAAGGTCCCC TGAAAAGGTCCCC TGAAAAGGTCCCC TGAAAAGGTCCCC TGAAAAGGTCCCCT TGAAAAGGTCCCT TGAAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAAAAAAA	260 STGAGCCGCAY STGAGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCACCGT 400 GGACCGT	TCACTICCACAC 270 TCACTICCACAC TCACTICCACACAC TCACTICCACACACACACACACACACACACACACACACAC	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 195 PAGTOC 195 PATAT 350 PATAT 289 PATAT 289 PATAT 280 PATAT 280 PATAT 280 PATAT 265
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.2.3 seq 6.1.4 seq 1.2.3 seq 1.1.4 seq 1.2.3 seq	GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC GTGGCTCC TGATACTC TGATACTC TGATACTC TGATACTC TGATACTC TGATACTC TGATACTC TGATACTC TGATACTC TGTAAATAI TGTAAATAI TGTAAATAI	ATCTTACK ATTTACT ATCTTACK AT	230 CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA 300 CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA CCAGATCA CATGTACA CCAGATCA	CGCTAGCTG 240 CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG TGTCCCCCGG TGTCCCCCGG TGTCCCCGGG TGTCCCCGGG TGTCCCCGGG TAGTGTATATA 380 TAGTGTATATA TAGTGTATATA	TGAAAGGTCCC 250 TGAAAGGTCCCC TCGTCGGCGTC TCGT	260 STGAGCCGCAY 260 STGAGCCGCAY STGAGCAGCTGAG AGCAGCTGAG AGCACGT AGG	TCACTICCACAC 270 TCACTICCACAC TCACTICCACACAC TCACTICCACACACACACACACACACACACACACACACAC	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 TATAT 350 TATAT 289 TATAT 289 TATAT 280 TATAT 265 TATAT 265
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.2.3 seq 6.1.4 seq	GTGGCTCC GTGATACTG TGATACTG TGATACTG TGTAAATAI TGTAAATAI TGTAAATAI TGTAAATAI	ATTTAATC ATTTAATC ATTTAAC ATCTTAC ATTTAATC ATTT	230 CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA 300 CCAGATCA CCAGATCA CCATGTACA CATGTACA CCATGTACA CCATGTA	CGCTAGCTG 240 CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG TGTCCCCCGG TGTCCCCCGGC TGTCCCCGGC TGTCCCGC TGTCCCCGGC TGTCCCCGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGCGC TGTCCCCGC TGTCCCCGCGC TGTCCCCGC TGTCCCCGC TGTCCCCGC TGTCCCCCGC TGTCCCCCGC TGTCCCCCGC TGTCCCCCGC	TGAAAGGTCCC 250 TGAAAGGTCCCC TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAAGGTCCCT TGAAAAAGGTCCCT TGAAAAAAAAAA	Zéo TGAGCCGCAT Zéo TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCTAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCACCGT GGACCGT GGACCGT GGACCGT	TCACTICCACAC 270 TCACTICCACAC TCACTICCACACAC TCACTICCACACACACACACACACACACACACACACACAC	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 195 PAGTOC 195 PATAT 350 PATAT 289 PATAT 289 PATAT 280 PATAT 280 PATAT 280 PATAT 265
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.2.3 seq 6.1.4 seq 1.2.3 seq 1.1.4 seq 1.2.3 seq	GTGGCTCC GTGATACTG TGATACTG TGATACTG TGTAAATAI TGTAAATAI TGTAAATAI TGTAAATAI	ATTTAATC ATTTAATC ATTTAAC ATCTTAC ATTTAATC ATTT	230 CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA 300 CCAGATCA CCAGATCA CCATGTACA CATGTACA CCATGTACA CCATGTA	CGCTAGCTG 240 CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG TGTCCCCCGG TGTCCCCCGGC TGTCCCCGGC TGTCCCGC TGTCCCCGGC TGTCCCCGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGCGC TGTCCCCGC TGTCCCCGCGC TGTCCCCGC TGTCCCCGC TGTCCCCGC TGTCCCCCGC TGTCCCCCGC TGTCCCCCGC TGTCCCCCGC	TGAAAGGTCCC 250 TGAAAGGTCCCC TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAAGGTCCCT TGAAAAAGGTCCCT TGAAAAAAAAAA	Zéo TGAGCCGCAT Zéo TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCTAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCACCGT GGACCGT GGACCGT GGACCGT	TGACTGCAGAC 270 TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC ACAAAATGTA ACAAAATGTA ACAAAATGTA	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 TATAT 350 TATAT 289 TATAT 289 TATAT 280 TATAT 265 402 341
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.2.3 seq 6.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.2 seq 6.2.2 seq 6.2.2 seq	GTGGCTCC GTGATACTG TGATACTG TGATACTG TGTAAATAI TGTAAATAI TGTAAATAI TGTAAATAI	ATTTAATC ATTTAATC ATTTAAC ATCTTAC ATTTAATC ATTT	230 CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA 300 CCAGATCA CCAGATCA CCATGTACA CATGTACA CCATGTACA CCATGTA	CGCTAGCTG 240 CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG TGTCCCCCGG TGTCCCCCGGC TGTCCCCGGC TGTCCCGC TGTCCCCGGC TGTCCCCGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGGC TGTCCCCGCGC TGTCCCCGC TGTCCCCGCGC TGTCCCCGC TGTCCCCGC TGTCCCCGC TGTCCCCCGC TGTCCCCCGC TGTCCCCCGC TGTCCCCCGC	TGAAAGGTCCC 250 TGAAAGGTCCCC TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAAGGTCCCT TGAAAAAGGTCCCT TGAAAAAAAAAA	Zéo TGAGCCGCAT Zéo TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCTAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCACCGT GGACCGT GGACCGT GGACCGT	TGACTGCAGAC 270 TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC ACAAAATGTA ACAAAATGTA ACAAAATGTA	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 PATAT 350 PATAT 289 PATAT 289 PATAT 280 PATAT 265 402 341 334 332
1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.4 seq 1.2.3 seq 6.1.4 seq 1.2.3 seq 6.2.2 seq 6.1.2 seq 6.2.2 seq 6.2.2 seq	GTGGCTCC GTGATACTG TGATACTG TGATACTG TGTAAATAI TGTAAATAI TGTAAATAI TGTAAATAI	ATTTAATC ATTTAATC ATTTAAC ATCTTAC ATTTAATC ATTT	230 CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA CCTAGTCA 300 CCAGATCA CCAGATCA CCATGTACA CATGTACA CCATGTACA CCATGTA	CGCTAGCTG 240 CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG CGCTAGCTG TGTCCCCCGG TGTCCCCCGG TGTCCCCGGG TGTCCCCGGG TGTCCCCGGG TAGTGTATATA 380 TAGTGTATATA TAGTGTATATA	TGAAAGGTCCC 250 TGAAAGGTCCCC TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAGGTCCCCT TGAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAGGTCCCCT TGAAAAAGGTCCCCT TGAAAAAGGTCCCT TGAAAAAGGTCCCT TGAAAAAAAAAA	Zéo TGAGCCGCAT Zéo TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCAT TGAGCCGCTAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCAGCTGAG AGCACCGT GGACCGT GGACCGT GGACCGT	TGACTGCAGAC 270 TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC TGACTGCAGAC ACAAAATGTA ACAAAATGTA ACAAAATGTA	280 PAGTOC 280 PAGTOC 219 PAGTOC 212 PAGTOC 210 PAGTOC 195 TATAT 350 TATAT 289 TATAT 289 TATAT 280 TATAT 265 402 341 334

FIGURE 20

HCV/BVDV chimera

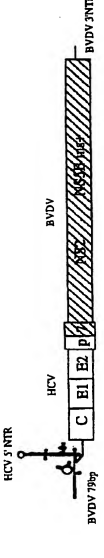


FIGURE 21

PCT/US99/08850

Gtatacgagaattagaaaaggcactcgtatacgtattgggcaattaaaaataattaggcctaggtacatggcacgtgccagcccct gatggggggacactccaccatgaatcactcccctgtgaggaactactgtcttcacgcagaaagcgtctagccatggcgttagtatgag tgtcgtgcagcctccaggacccccctcccgggagagccatagtggtctgcggaaccggtgagtacaccggaattgccaggacgac cgggtcctttcttggataaacccgctcaatgcctggagatttgggcgtgccccgcaagactgctagccgagtagtgttgggtcgcgaa AAGTTCCCGGGTGGCGGTCAGATCGTTGGTGGAGTTTACTTGTTGCCGCGCAG GGGCCCTAGATTGGGTGTGCGCGCGACGAGGAAGACTTCCGAGCGGTCGCAA CCTCGAGGTAGACGTCAGCCTATCCCCAAGGCACGTCGGCCCGAGGGCAGGA CCTGGGCTCAGCCCGGGTACCCTTGGCCCCTCTATGGCAATGAGGGTTGCGGG TGGGCGGGATGGCTCCTGTCTCCCCGTGGCTCTCGGCCTAGCTGGGGCCCCAC AGACCCCGGCGTAGGTCGCGCAATTTGGGTAAGGTCATCGATACCCTTACGT GCGGCTTCGCCGACCTCATGGGGTACATACCGCTCGTCGGCGCCCCTCTTGGA GGCGCTGCCAGGGCCCTGGCGCATGGCGTCCGGGTTCTGGAAGACGGCGTGA ACTATGCAACAGGGAACCITCCTGGTTGCTCTTTCTCTATCTTCCTTCTGGCCCT GCTCTCTTGCCTGACCGTGCCCGCTTCAGCCTACCAAGTGCGCAATTCCTCGGG GCTTTACCATGTCACCAATGATTGCCCTAACTCGAGTATTGTGTACGAGGCGGC CGATGCCATCCTGCACACTCCGGGGTGTGTCCCTTGCGTTCGCGAGGGTAACG CCTCGAGGTGTTGGGTGGCGGTGACCCCCACGGTGGCCACCAGGGACGGCAA ACTCCCCACAACGCAGCTTCGACGTCATATCGATCTGCTTGTCGGGAGCGCCA GTCAACTGTTTACCTTCTCCCAGGCGCCACTGGACGACGCAAGACTGCAATT GTTCTATCTATCCCGGCCATATAACGGGTCATCGCATGGCATGGGATATGATGA TGAACTGGTCCCCTACGGCAGCGTTGGTGGTAGCTCAGCTGCTCCGGATCCCA CAAGCCATCATGGACATGATCGCTGGTGCTCACTGGGGAGTCCTGGCGGGCAT AGCGTATTTCTCCATGGTGGGGAACTGGGCGAAGGTCCTGGTAGTGCTGCTGC CACCACGCTGGGCTTGTTGGTCTCCTTACACCAGGCGCCCAAGCAGAACATCC AACTGATCAACACCAACGGCAGTTGGCACATCAATAGCACGGCCTTGAACTGC AATGAAAGCCTTAACACCGGCTGGTTAGCAGGGCTCTTCTATCAGCACAAATTC AACTCTTCAGGCTGTCCTGAGAGGTTGGCCAGCTGCCGACGCCTTACCGATTTT GCCCAGGGCTGGGGTCCTATCAGTTATGCCAACGGAAGCGGCCTCGACGAAC GCCCTACTGCTGGCACTACCCTCCAAGACCTTGTGGCATTGTGCCCGCAAAG AGCGTGTGTGGCCCGGTATATTGCTTCACTCCCAGCCCCGTGGTGGTGGGAAC GACCGACAGGTCGGGCGCCCTACCTACAGCTGGGGTGCAAATGATACGGAT GTCTTCGTCCTTAACAACACCAGGCCACCGCTGGGCAATTGGTTCGGTTGTACC TGGATGAACTCAACTGGATTCACCAAAGTGTGCGGAGCGCCCCCTTGTGTCAT CGGAGGGTGGCAACACACCTTGCTCTGCCCCACTGATTGTTTCCGCAAGC ATCCGGAAGCCACATACTCTCGGTGCGGCTCCGGTCCCTGGATTACACCCAGG TGCATGGTCGACTACCCGTATAGGCTTTGGCACTATCCTTGTACCATCAATTAC ACCATATTCAAAGTCAGGATGTACGTGGGAGGGGTCGAGCACAGGCTGGAAG CGGCCTGCAACTGGACGCGGGGCGAACGCTGTGATCTGGAAGACAGGGACAG GTCCGAGCTCAGCCCATTGCTGCTGTCCACCACAGTGGCAGGTCCTTCCGT GTTCTTTCACGACCCTGCCAGCCTTGTCCACCGGCCTCATCCACCTCCACCAGA ACATTGTGGACGTGCAGTACTTGTACGGGGTAGGGTCAAGCATCGCGTCCTGG GCCATTAAGTGGGAGTACGTCGTTCTCCTGTTCCTGCTTGCAGACGCGCGC GTCTGCTCCTGCTTGTGGATGATGTTACTCATATCCCAAGCGGAGGCGGCTTTG GAGAACCTCGTAATACTCAATGCAGCATCCCTGGCCGGGACGCACGGTCTTGT

CGGAGCGGTCTACGCCTTCTACGGGAAGTGGGTCTTACTCTTATACCACATCTT AGTGGTACACCCAATCAAATCTGTAATTGTGATCCTACTGATGATTGGGGATGT GGTAAAGGCCGATTCAGGGGGCCAAGAGTACTTGGGGAAAATAGACCTCTGTT TTACAACAGTAGTACTAATCGTCATAGGTTTAATCATAGCTAGGCGTGACCCAA CTATAGTGCCACTGGTAACAATAATGGCAGCACTGAGGGTCACTGAACTGACC CACCAGCCTGGAGTTGACATCGCTGTGGCGGTCATGACTATAACCCTACTGAT GGTTAGCTATGTGACAGATTATTTTAGATATAAAAAATGGTTACAGTGCATTCTC AGCCTGGTATCTGCGGTGTTCTTGATAAGAAGCCTAATATACCTAGGTAGAATC GAGATGCCAGAGGTAACTATCCCAAACTGGAGACCACTAACTTTAATACTATTA TTGTTGCAATGTGTGCCTATCTTATTGCTGGTCACAACCTTGTGGGCCGACTTCT TAACCCTAATACTGATCCTGCCTACCTATGAATTGGTTAAATTATACTATCTGAA **AACTGTTAGGACTGATACAGAAAGAAGTTGGCTAGGGGGGATAGACTATACAA** GAGTTGACTCCATCTACGACGTTGATGAGAGTGGAGAGGGCGTATATCTTTTTC CATCAAGGCAGAAAGCACAGGGGAATTTTTCTATACTCTTGCCCCTTATCAAAG CAACACTGATAAGTTGCGTCAGCAGTAAATGGCAGCTAATATACATGAGTTACT TAACTTTGGACTTTATGTACTACATGCACAGGAAAGTTATAGAAGAGATCTCAG GAGGTACCAACATAATATCCAGGTTAGTGGCAGCACTCATAGAGCTGAACTGG TCCATGGAAGAAGAGGAGAGCAAAGGCTTAAAGAAGTTTTATCTATTGTCTGG AAGGTTGAGAAACCTAATAATAAAACATAAGGTAAGGAATGAGACCGTGGCTT CTTGGTACGGGGGGGGGGAGTCTACGGTATGCCAAAGATCATGACTATAATC AAGGCCAGTACACTGAGTAAGAGCAGGCACTGCATAATATGCACTGTATGTGA GGGCCGAGAGTGGAAAGGTGGCACCTGCCCAAAATGTGGACGCCATGGGAAG **AATCTTTATAAGGGAAGGCAACTTTGAGGGTATGTGCAGCCGATGCCAGGGAA** AGCATAGGAGGTTTGAAATGGACCGGGAACCTAAGAGTGCCAGATACTGTGCT GAGTGTAATAGGCTGCATCCTGCTGAGGAAGGTGACTTTTGGGCAGAGTCGAG TATCACAGAGTGGGCTGGATGCCAGCGTGTGGGAATCTCCCCAGATACCCACA ATGGCTTTGTACAATATACCGCTAGGGGGCAACTATTTCTGAGAAACTTGCCCG TACTGGCAACTAAAGTAAAAATGCTCATGGTAGGCAACCTTGGAGAAGAAATT GGTAATCTGGAACATCTTGGGTGGATCCTAAGGGGGCCTGCCGTGTGTAAGAA GATCACAGAGCACGAAAAATGCCACATTAATATACTGGATAAACTAACCGCATT TTTCGGGATCATGCCAAGGGGGACTACACCCAGAGCCCCGGTGAGGTTCCCTA CAAGGCGGGATAAGTTCAGTCGACCATGTAACCGCCGGAAAAGATCTACTGGT CTGTGACAGCATGGGACGAACTAGAGTGGTTTGCCAAAGCAACAACAGGTTGA CCGATGAGACAGAGTATGGCGTCAAGACTGACTCAGGGTGCCCAGACGGTGC CAGATGTTATGTGTTAAATCCAGAGGCCGTTAACATATCAGGATCCAAAGGGG CAGTCGTTCACCTCCAAAAGACAGGTGGAGAATTCACGTGTGTCACCGCATCA GGCACACCGGCTTTCTTCGACCTAAAAAACTTGAAAGGATGGTCAGGCTTGCCT ATATTTGAAGCCTCCAGCGGGAGGGTGGTTGGCAGAGTCAAAGTAGGGAAGA ATGAAGAGTCTAAACCTACAAAAATAATGAGTGGAATCCAGACCGTCTCAAAAA ACAGAGCAGACCTGACCGAGATGGTCAAGAAGATAACCAGCATGAACAGGGG AGACTTCAAGCAGATTACTTTGGCAACAGGGGCAGGCAAAACCACAGAACTCC CAAAAGCAGTTATAGAGGAGATAGGAAGACACAAGAGAGTATTAGTTCTTATA CCATTAAGGGCAGCGGCAGAGTCAGTCTACCAGTATATGAGATTGAAACACCC AAGCATCTCTTTTAACCTAAGGATAGGGGACATGAAAGAGGGGGACATGGCAA CCGGGATAACCTATGCATCATACGGGTACTTCTGCCAAATGCCTCAACCAAAGC TCAGAGCTGCTATGGTAGAATACTCATACATATTCTTAGATGAATACCATTGTGC CACTCCTGAACAACTGGCAATTATCGGGAAGATCCACAGATTTTCAGAGAGTAT AAGGGTTGTCGCCATGACTGCCACGCCAGCAGGGTCGGTGACCACAACAGGT CAAAAGCACCCAATAGAGGAATTCATAGCCCCCGAGGTAATGAAAGGGGAGG

ATCTTGGTAGTCAGTTCCTTGATATAGCAGGGTTAAAAATACCAGTGGATGAGA TGAAAGGCAATATGTTGGTTTTTGTACCAACGAGAAACATGGCAGTAGAGGTA GCAAAGAAGCTAAAAGCTAAGGGCTATAACTCTGGATACTATTACAGTGGAGA GGATCCAGCCAATCTGAGAGTTGTGACATCACAATCCCCCTATGTAATCGTGGC TACAAATGCTATTGAATCAGGAGTGACACTACCAGATTTGGACACGGTTATAGA CACGGGGTTGAAATGTGAAAAGAGGGTGAGGGTATCATCAAAGATACCCTTCA TCGTAACAGGCCTTAAGAGGATGGCCGTGACTGTGGGTGAGCAGGCGCAGCG TAGGGGCAGAGTAGGTGAAACCCGGGAGGTATTATAGGAGCCAGGAA ACAGCAACAGGGTCAAAGGACTACCACTATGACCTCTTGCAGGCACAAAGATA CGGGATTGAGGATGGAATCAACGTGACGAAATCCTTTAGGGAGATGAATTACG ATTGGAGCCTATACGAGGAGGACAGCCTACTAATAACCCAGCTGGAAATACTA AATAATCTACTCATCTCAGAAGACTTGCCAGCCGCTGTTAAGAACATAATGGCC AGGACTGATCACCCAGAGCCAATCCAACTTGCATACAACAGCTATGAAGTCCA GGTCCCGGTCCTATTCCCAAAAATAAGGAATGGAGAAGTCACAGACACCTACG AAAATTACTCGTTTCTAAATGCCAGAAAGTTAGGGGAGGATGTGCCCGTGTATA TCTACGCTACTGAAGATGAGGATCTGGCAGTTGACCTCTTAGGGCTAGACTGG CCTGATCCTGGGAACCAGCAGGTAGTGGAGACTGGTAAAGCACTGAAGCAAGT GACCGGGTTGTCCTCGGCTGAAAATGCCCTACTAGTGGCTTTATTTGGGTATGT GGGTTACCAGGCTCTCTCAAAGAGGCATGTCCCAATGATAACAGACATATATAC CATCGAGGACCAGAGACTAGAAGACACCACCCACCTCCAGTATGCACCCAACG CCATAAAAACCGATGGGACAGAGACTGAACTGAAAGAACTGGCGTCGGGTGA CGTGGAAAAAATCATGGGAGCCATTTCAGATTATGCAGCTGGGGGACTGGAGT TTGTTAAATCCCAAGCAGAAAAGATAAAAACAGCTCCTTTGTTTAAAGAAAACG CAGAAGCCGCAAAAGGGTATGTCCAAAAATTCATTGACTCATTAATTGAAAATA AGCATAGCTGCAAGACTGGGGCATGAAACAGCGTTTGCCACACTAGTGTTAAA **GTGGCTAGCTTTTGGAGGGGAATCAGTGTCAGACCACGTCAAGCAGGCGGCA** GAGACACAGCAAGAAGGGAGGCGATTCGTCGCAAGCCTGTTCATCTCCGCACT GGCAACCTACACATACAAAACTTGGAATTACCACAATCTCTCTAAAGTGGTGGA ACCAGCCCTGGCTTACCTCCCCTATGCTACCAGCGCATTAAAAATGTTCACCCC AACGCGGCTGGAGAGCGTGGTGATACTGAGCACCACGATATATAAAACATACC TCTCTATAAGGAAGGGGAAGAGTGATGGATTGCTGGGTACGGGGATAAGTGC GGGGGTAGGGCAATCGCTGCGCACAACGCTATTGAGTCCAGTGAACAGAAA AGGACCCTACTTATGAAGGTGTTTGTAAAGAACTTCTTGGATCAGGCTGCAACA GATGAGCTGGTAAAAGAAAACCCAGAAAAAATTATAATGGCCTTATTTGAAGCA GTCCAGACAATTGGTAACCCCCTGAGACTAATATACCACCTGTATGGGGTTTAC TATTCACATTGATAATGTTTGAAGCCTTCGAGTTATTAGGGATGGACTCACAAG GGAAAATAAGGAACCTGTCCGGAAATTACATTTTGGATTTGATATACGGCCTAC ACAAGCAAATCAACAGAGGGCTGAAGAAAATGGTACTGGGGTGGGCCCCTGC ACCCTTTAGTTGTGACTGGACCCCTAGTGACGAGGGGATCAGATTGCCAACAG ACAACTATTTGAGGGTAGAAACCAGGTGCCCATGTGGCTATGAGATGAAAGCT TTCAAAAATGTAGGTGGCAAACTTACCAAAGTGGAGGAGAGCGGGCCTTTCCT ATGTAGAAACAGACCTGGTAGGGGACCAGTCAACTACAGAGTCACCAAGTATT ACGATGACAACCTCAGAGAGATAAAACCAGTAGCAAAGTTGGAAGGACAGGTA GAGCACTACTACAAAGGGGTCACAGCAAAAATTGACTACAGTAAAGGAAAAAT GCTCTTGGCCACTGACAAGTGGGAGGTGGAACATGGTGTCATAACCAGGTTAG CTAAGAGATATACTGGGGTCGGGTTCAATGGTGCATACTTAGGTGACGAGCCC AATCACCGTGCTCTAGTGGAGAGGGACTGTGCAACTATAACCAAAAACACAGT ACAGTTTCTAAAAATGAAGAAGGGGTGTGCGTTCACCTATGACCTGACCATCTC CAATCTGACCAGGCTCATCGAACTAGTACACAGGAACAATCTTGAAGAGAAGG

ACGTAGGGACTATAAAACCAGTACTAGGAGAGAGAGTAATCCCCGACCCTGTA GTTGATATCAATTTACAACCAGAGGTGCAAGTGGACACGTCAGAGGTTGGGAT CACAATAATTGGAAGGGAAACCCTGATGACAACGGGAGTGACACCTGTCTTGG AAAAAGTAGAGCCTGACGCCAGCGACAACCAAAACTCGGTGAAGATCGGGTTG GATGAGGGTAATTACCCAGGGCCTGGAATACAGACACATACACTAACAGAAGA AATACACAACAGGGATGCGAGGCCCTTCATCATGATCCTGGGCTCAAGGAATT CCATATCAAATAGGGCAAAGACTGCTAGAAATATAAATCTGTACACAGGAAATG ACCCCAGGGAAATACGAGACTTGATGGCTGCAGGGCGCATGTTAGTAGTAGCA CTGAGGGATGTCGACCCTGAGCTGTCTGAAATGGTCGATTTCAAGGGGACTTT TTTAGATAGGGAGGCCCTGGAGGCTCTAAGTCTCGGGCAACCTAAACCGAAGC AGGTTACCAAGGAAGCTGTTAGGAATTTGATAGAACAGAAAAAAGATGTGGAG ATCCCTAACTGGTTTGCATCAGATGACCCAGTATTTCTGGAAGTGGCCTTAAAA **AATGATAAGTACTACTTAGTAGGAGATGTTGGAGAGCTAAAAGATCAAGCTAAA** GCACTTGGGGCCACGGATCAGACAAGAATTATAAAGGAGGTAGGCTCAAGGA GTTTAACTCCACTGTTTGAGGAATTGTTGCTACGGTGCCCACCTGCAACTAAGA GCAATAAGGGGCACATGGCATCAGCTTACCAATTGGCACAGGGTAACTGGGAG ACACCCATATGAAGCTTACCTGAAGTTGAAAGATTTCATAGAAGAAGAAGAAGAA GAAACCTAGGGTTAAGGATACAGTAATAAGAGAGCACAACAAATGGATACTTA AAAAATAAGGTTTCAAGGAAACCTCAACACCAAGAAAATGCTCAACCCAGGG AAACTATCTGAACAGTTGGACAGGGAGGGGGGGCGCAAGAGGGAACATCTACAACCA CCAGATTGGTACTATAATGTCAAGTGCAGGCATAAGGCTGGAGAAATTGCCAA TAGTGAGGGCCCAAACCGACACCAAAACCTTTCATGAGGCAATAAGAGATAAG ATAGACAAGAGTGAAAACCGGCAAAATCCAGAATTGCACAACAAATTGTTGGA GATTTTCCACACGATAGCCCAACCCACCCTGAAACACACCTACGGTGAGGTGA CGTGGGAGCAACTTGAGGCGGGGGTAAATAGAAAGGGGGCAGCAGGCTTCCT GGAGAAGAAGAACATCGGAGAAGTATTGGATTCAGAAAAGCACCTGGTAGAAC AATTGGTCAGGGATCTGAAGGCCGGGAGAAAGATAAAATATTATGAAACTGCA TGGTGGTTGAGAAGAGGCCAAGAGTTATCCAATACCCTGAAGCCAAGACAAGG CTAGCCATCACTAAGGTCATGTATAACTGGGTGAAACAGCAGCCCGTTGTGATT CCAGGATATGAAGGAAAGACCCCCTTGTTCAACATCTTTGATAAAGTGAGAAAG GAATGGGACTCGTTCAATGAGCCAGTGGCCGTAAGTTTTGACACCAAAGCCTG GGACACTCAAGTGACTAGTAAGGATCTGCAACTTATTGGAGAAATCCAGAAATA TTACTATAAGAAGGAGTGGCACAAGTTCATTGACACCATCACCGACCACATGAC GAGGGAGCGGCCAGCCAGACACAGTGCTGGCAACAGCATGTTAAATGTCCT GACAATGATGTACGGCTTCTGCGAAAGCACAGGGGTACCGTACAAGAGTTTCA ACAGGGTGGCAAGGATCCACGTCTGTGGGGATGATGGCTTCTTAATAACTGAA **AAAGGGTTAGGGCTGAAATTTGCTAACAAAGGGATGCAGATTCTTCATGAAGC** AGGCAAACCTCAGAAGATAACGGAAGGGGAAAAGATGAAAGTTGCCTATAGAT TTGAGGATATAGAGTTCTGTTCTCATACCCCAGTCCCTGTTAGGTGGTCCGACA ACACCAGTAGTCACATGGCCGGGAGAGACACCGCTGTGATACTATCAAAGATG GCAACAAGATTGGATTCAAGTGGAGAGAGGGGTACCACAGCATATGAAAAAGC GGTAGCCTTCAGTTTCTTGCTGATGTATTCCTGGAACCCGCTTGTTAGGAGGAT TTGCCTGTTGGTCCTTTCGCAACAGCCAGAGACAGACCCATCAAAACATGCCAC TTATTATTACAAAGGTGATCCAATAGGGGCCTATAAAGATGTAATAGGTCGGAA TCTAAGTGAACTGAAGAGAACAGGCTTTGAGAAATTGGCAAATCTAAACCTAAG CCTGTCCACGTTGGGGGTCTGGACTAAGCACACAAGCAAAAGAATAATTCAGG ACTGTGTTGCCATTGGGAAAGAAGAGGGCAACTGGCTAGTTAAGCCCGACAGG CTGATATCCAGCAAAACTGGCCACTTATACATACCTGATAAAGGCTTTACATTAC AAGGAAAGCATTATGAGCAACTGCAGCTAAGAACAGAGACAAACCCGGTCATG GGGGTTGGGACTGAGAGATACAAGTTAGGTCCCATAGTCAATCTGCTGCTGAG

AAGGTTGAAAATTCTGCTCATGACGGCCGTCGGCGTCAGCAGCTGAgacaaaatgtat atattgtaaataaattaatccatgtacatagtgtatataaattaagttgggaccgtccacctcaagaagacgacacgcccaacacgcacag ctaaacagtagtcaagattatctacctcaagataacactacatttaatgcacacagcactttagctgtatgaggatacgcccgacgtctatag ttggactaggggaagacctctaacagccccc

IGURE 22-5

HCV/BVDV chimera with selectable marker



FIGURE 23

Gtatacgagaattagaaaaggcactcgtatacgtattgggcaattaaaaataataattaggcctaggtacatggcacgtgccagcccct galgggggcgacactccaccatgaatcactcccctgtgaggaactactgtcttcacgcagaaagcgtctagccatggcgttagtatgag tgtcgtgcagcctccaggacccccctcccgggagagccatagtggtctgcggaaccggtgagtacaccggaattgccaggacgac cgggtcctttcttggataaacccgctcaatgcctggagatttgggcgtgcccccgcaagactgctagccgagtagtgttgggtcgcgaa aggccttgtggtactgcctgatagggtgcttgcgagtgccccgggaggtctcgtagaccgtgcaccATGAGCACGAATCCTAAACCTCAAAGAAAAACCAAACGTAACACCAACCGTCGCCCACAGGACGTC AAGTTCCCGGGTGGCGGTCAGATCGTTGGTGGAGTTTACTTGTTGCCGCGCAG GGGCCCTAGATTGGGTGTGCGCGCGACGAGGAAGACTTCCGAGCGGTCGCAA CCTCGAGGTAGACGTCAGCCTATCCCCAAGGCACGTCGGCCCGAGGGCAGGA CCTGGGCTCAGCCCGGGTACCCTTGGCCCCTCTATGGCAATGAGGGTTGCGGG TGGGCGGATGCTCCTGTCTCCCCGTGGCTCTCGGCCTAGCTGGGGCCCCAC AGACCCCGGCGTAGGTCGCGCAATTTGGGTAAGGTCATCGATACCCTTACGT GCGGCTTCGCCGACCTCATGGGGTACATACCGCTCGTCGGCGCCCCTCTTGGA GGCGCTGCCAGGGCCCTGGCGCATGGCGTCCGGGTTCTGGAAGACGGCGTGA GCTTTACCATGTCACCAATGATTGCCCTAACTCGAGTATTGTGTACGAGGCGGC CGATGCCATCCTGCACACTCCGGGGTGTGTCCCTTGCGTTCGCGAGGGTAACG CCTCGAGGTGTTGGGTGGCGGTGACCCCCACGGTGGCCACCAGGGACGGCAA ACTCCCCACAACGCAGCTTCGACGTCATATCGATCTGCTTGTCGGGAGCGCCA GTCAACTGTTTACCTTCTCCCAGGCGCCACTGGACGACGCAAGACTGCAATT GTTCTATCTATCCCGGCCATATAACGGGTCATCGCATGGCATGGGATATGATGA TGAACTGGTCCCCTACGGCAGCGTTGGTGGTAGCTCAGCTGCTCCGGATCCCA CAAGCCATCATGGACATGATCGCTGGTGCTCACTGGGGAGTCCTGGCGGCAT AGCGTATTTCTCCATGGTGGGGAACTGGGCGAAGGTCCTGGTAGTGCTGCTGC CACCACGGCTGGGCTTGTTGGTCTCCTTACACCAGGCGCCAAGCAGAACATCC AACTGATCAACACCAACGGCAGTTGGCACATCAATAGCACGGCCTTGAACTGC AATGAAAGCCTTAACACCGGCTGGTTAGCAGGGCTCTTCTATCAGCACAAATTC AACTCTTCAGGCTGTCCTGAGAGGTTGGCCAGCTGCCGACGCCTTACCGATTTT GCCCAGGGCTGGGGTCCTATCAGTTATGCCAACGGAAGCGGCCTCGACGAAC GCCCTACTGCTGGCACTACCCTCCAAGACCTTGTGGCATTGTGCCCGCAAAG AGCGTGTGTGGCCCGGTATATTGCTTCACTCCCAGCCCCGTGGTGGTGGGAAC GACCGACAGGTCGGGCGCCCTACCTACAGCTGGGGTGCAAATGATACGGAT GTCTTCGTCCTTAACAACACCAGGCCACCGCTGGGCAATTGGTTCGGTTGTACC TGGATGAACTCAACTGGATTCACCAAAGTGTGCGGAGCGCCCCCTTGTGTCAT CGGAGGGGTGGCCAACACACCTTGCTCTGCCCCACTGATTGTTTCCGCAAGC ATCCGGAAGCCACATACTCTCGGTGCGGCTCCGGTCCCTGGATTACACCCAGG TGCATGGTCGACTACCCGTATAGGCTTTGGCACTATCCTTGTACCATCAATTAC ACCATATTCAAAGTCAGGATGTACGTGGGAGGGGTCGAGCACAGGCTGGAAG CGGCCTGCAACTGGACGCGGGGCGAACGCTGTGATCTGGAAGACAGGGACAG GTCCGAGCTCAGCCCATTGCTGCTGTCCACCACAGTGGCAGGTCCTTCCGT GTTCTTCACGACCCTGCUAGCCTTGTCCACCGGCCTCATCCACCTCCACCAGA ACATTGTGGACGTGCAGTACTTGTACGGGGTAGGGTCAAGCATCGCGTCCTGG GCCATTAAGTGGGAGTACGTCGTTCTCCTGTTCCTCCTGCTTGCAGACGCGCGC GTCTGCTCCTGCTTGTGGATGATGTTACTCATATCCCAAGCGGAGGCGGCTTTG GAGAACCTCGTAATACTCAATGCAGCATCCCTGGCCGGGACGCACGGTCTTGT

CGGAGCGGTCTACGCCTTCTACGGGAAGTGGGTCTTACTCTTATACCACATCTT AGTGGTACACCCAATCAAATCTGTAATTGTGATCCTACTGATGATTGGGGATGT GGTAAAGGCCGATTCAGGGGGCCAAGAGTACTTGGGGAAAATAGACCTCTGTT TTACAACAGTAGTACTAATCGTCATAGGTTTAATCATAGCTAGGCGTGACCCAA CTATAGTGCCACTGGTAACAATAATGGCAGCACTGAGGGTCACTGAACTGACC CACCAGCCTGGAGTTGACATCGCTGTGGCGGTCATGACTATAACCCTACTGAT GGTTAGCTATGTGACAGATTATTTTAGATATAAAAAATGGTTACAGTGCATTCTC AGCCTGGTATCTGCGGTGTTCTTGATAAGAAGCCTAATATACCTAGGTAGAATC GAGATGCCAGAGGTAACTATCCCAAACTGGAGACCACTAACTTTAATACTATTA TTGTTGCAATGTGTGCCTATCTTATTGCTGGTCACAACCTTGTGGGCCGACTTCT TAACCCTAATACTGATCCTGCCTACCTATGAATTGGTTAAATTATACTATCTGAA **AACTGTTAGGACTGATACAGAAAGAAGTTGGCTAGGGGGGATAGACTATACAA** GAGTTGACTCCATCTACGACGTTGATGAGAGTGGAGAGGGCGTATATCTTTTTC CATCAAGGCAGAAAGCACAGGGGAATTTTTCTATACTCTTGCCCCTTATCAAAG CAACACTGATAAGTTGCGTCAGCAGTAAATGGCAGCTAATATACATGAGTTACT TAACITTGGACITTATGTACTACATGCACAGGAAAGTTATAGAAGAGATCTCAG GAGGTACCAACATAATATCCAGGTTAGTGGCAGCACTCATAGAGCTGAACTGG TCCATGGAAGAAGAGGAGAGCAAAGGCTTAAAGAAGTTTTATCTATTGTCTGG AAGGTTGAGAAACCTAATAATAAAACATAAGGTAAGGAATGAGACCGTGGCTT CTTGGTACGGGGAGGAGGAAGTCTACGGTATGCCAAAGATCATGACTATAATC AAGGCCAGTACACTGAGTAAGAGCAGGCACTGCATAATATGCACTGTATGTGA GGGCCGAGAGTGGAAAGGTGGCACCTGCCCAAAATGTGGACGCCATGGGAAG **AATCTTTATAAGGGAAGGCAACTTTGAGGGTATGTGCAGCCGATGCCAGGGAA** AGCATAGGAGGTTTGAAATGGACCGGGAACCTAAGAGTGCCAGATACTGTGCT GAGTGTAATAGGCTGCATCCTGCTGAGGAAGGTGACTTTTGGGCAGAGTCGAG TATCACAGAGTGGGCTGGATGCCAGCGTGTGGGAATCTCCCCAGATACCCACA ATGGCTTTGTACAATATACCGCTAGGGGGCAACTATTTCTGAGAAACTTGCCCG TACTGGCAACTAAAGTAAAAATGCTCATGGTAGGCAACCTTGGAGAAGAAATT GGTAATCTGGAACATCTTGGGTGGATCCTAAGGGGGCCTGCCGTGTGTAAGAA GATCACAGAGCACGAAAAATGCCACATTAATATACTGGATAAACTAACCGCATT TTTCGGGATCATGCCAAGGGGGACTACACCCAGAGCCCCGGTGAGGTTCCCTA CAAGGCGGGATAAGTTCAGTCGACCATGTAACCGCCGGAAAAGATCTACTGGT CTGTGACAGCATGGGACGAACTAGAGTGGTTTGCCAAAGCAACAACAGGTTGA CCGATGAGACAGAGTATGGCGTCAAGACTGACTCAGGGTGCCCAGACGGTGC CAGATGTTATGTGTTAAATCCAGAGGCCGTTAACATATCAGGATCCAAAGGGG CAGTCGTTCACCTCCAAAAGACAGGTGGAGAATTCACGTGTGTCACCGCATCA GGCACACCGGCTTTCTTCGACCTAAAAAACTTGAAAGGATGGTCAGGCTTGCCT **ATATTTGAAGCCTCCAGCGGGAGGGTGGTTGGCAGAGTCAAAGTAGGGAAGA ATGAAGAGTCTAAACCTACAAAAATAATGAGTGGAATCCAGACCGTCTCAAAAA** ACAGAGCAGACCTGACCGAGATGGTCAAGAAGATAACCAGCATGAACAGGGG AGACTTCAAGCAGATTACTTTGGCAACAGGGGCAGGCAAAACCACAGAACTCC CAAAAGCAGTTATAGAGGAGATAGGAAGACACAAGAGAGTATTAGTTCTTATA CCATTAAGGGCAGCGCAGAGTCAGTCTACCAGTATATGAGATTGAAACACCC <u>AAGCATCTCTTTTAACCTAAGGATAGGGGACATGAAAGAGGGGGGACATGGCAA</u> CCGGGATAACCTATGCATCATACGGGTACTTCTGCCAAATGCCTCAACCAAAGC TCAGAGCTGCTATGGTAGAATACTCATACATATTCTTAGATGAATACCATTGTGC CACTCCTGAACAACTGGCAATTATCGGGAAGATCCACAGATTTTCAGAGAGTAT AAGGGTTGTCGCCATGACTGCCACGCCAGCAGGGTCGGTGACCACAACAGGT

CAAAAGCACCAATAGAGGAATTCATAGCCCCCGAGGTAATGAAAGGGGAGG **ATCTTGGTAGTCAGTTCCTTGATATAGCAGGGTTAAAAATACCAGTGGATGAGA** TGAAAGGCAATATGTTGGTTTTTGTACCAACGAGAAACATGGCAGTAGAGGTA GCAAAGAAGCTAAAAGCTAAGGGCTATAACTCTGGATACTATTACAGTGGAGA GGATCCAGCCAATCTGAGAGTTGTGACATCACAATCCCCCTATGTAATCGTGGC TACAAATGCTATTGAATCAGGAGTGACACTACCAGATTTGGACACGGTTATAGA CACGGGGTTGAAATGTGAAAAGAGGGTGAGGGTATCATCAAAGATACCCTTCA TCGTAACAGGCCTTAAGAGGATGGCCGTGACTGTGGGTGAGCAGGCGCAGCG TAGGGGCAGAGTAGGTAGAGTGAAACCCGGGAGGTATTATAGGAGCCAGGAA ACAGCAACAGGGTCAAAGGACTACCACTATGACCTCTTGCAGGCACAAAGATA CGGGATTGAGGATGGAATCAACGTGACGAAATCCTTTAGGGAGATGAATTACG ATTGGAGCCTATACGAGGAGGACAGCCTACTAATAACCCAGCTGGAAATACTA **AATAATCTACTCATCTCAGAAGACTTGCCAGCCGCTGTTAAGAACATAATGGCC** AGGACTGATCACCCAGAGCCAATCCAACTTGCATACAACAGCTATGAAGTCCA GGTCCCGGTCCTGTTCCCAAAAATAAGGAATGGAGAAGTCACAGACACCTACG AAAATTACTCGTTTCTAAATGCCAGAAAGTTAGGGGAGGATGTGCCCGTGTATA TCTACGCTACTGAAGATGAGGATCTGGCAGTTGACCTCTTAGGGCTAGACTGG CCTGATCCTGGGAACCAGCAGGTAGTGGAGACTGGTAAAGCACTGAAGCAAGT GACCGGGTTGTCCTCGGCTGAAAATGCCCTACTAGTGGCTTTATTTGGGTATGT GGGTTACCAGGCTCTCTCAAAGAGGCATGTCCCAATGATAACAGACATATATAC CATCGAGGACCAGAGACTAGAAGACACCACCCACCTCCAGTATGCACCCAACG CCATAAAAACCGATGGGACAGAGACTGAACTGAAAGAACTGGCGTCGGGTGA CGTGGAAAAAATCATGGGAGCCATTTCAGATTATGCAGCTGGGGGACTGGAGT TTGTTAAATCCCAAGCAGAAAAGATAAAAACAGCTCCTTTGTTTAAAGAAAACG CAGAAGCCGCAAAAGGGTATGTCCAAAAATTCATTGACTCATTAATTGAAAATA AGCATAGCTGCAAGACTGGGGCATGAAACAGCGTTTGCCACACTAGTGTTAAA GTGGCTAGCTTTTGGAGGGGAATCAGTGTCAGACCACGTCAAGCAGGCGGCA GAGACACAGCAAGAAGGGAGGCGATTCGTCGCAAGCCTGTTCATCTCCGCACT GGCAACCTACACATACAAAACTTGGAATTACCACAATCTCTCTAAAGTGGTGGA ACCAGCCTGGCTTACCTCCCTATGCTACCAGCGCATTAAAAATGTTCACCCC AACGCGGCTGGAGAGCGTGGTGATACTGAGCACCACGATATATAAAACATACC TCTCTATAAGGAAGGGGAAGAGTGATGGATTGCTGGGTACGGGGATAAGTGC GGGGTAGGGCAATCGCTGCGCACAACGCTATTGAGTCCAGTGAACAGAAA AGGACCTACTTATGAAGGTGTTTGTAAAGAACTTCTTGGATCAGGCTGCAACA GATGAGCTGGTAAAAGAAAACCCAGAAAAAATTATAATGGCCTTATTTGAAGCA GTCCAGACAATTGGTAACCCCCTGAGACTAATATACCACCTGTATGGGGTTTAC TATTCACATTGATAATGTTTGAAGCCTTCGAGTTATTAGGGATGGACTCACAAG GGAAAATAAGGAACCTGTCCGGAAATTACATTTTGGATTTGATATACGGCCTAC ACAAGCAAATCAACAGAGGGCTGAAGAAAATGGTACTGGGGTGGGCCCCTGC ACCCTTTAGTTGTGACTGGACCCCTAGTGACGAGGGATCAGATTGCCAACAG ACAACTATTTGAGGGTAGAAACCAGGTGCCCATGTGGCTATGAGATGAAAGCT TTCAAAAATGTAGGTGGCAAACTTACCAAAGTGGAGGAGAGCGGGCCTTTCCT ATGTAGAAACAGACCTGGTAGGGGACCAGTCAACTACAGAGTCACCAAGTATT ACGATGACAACCTCAGAGAGATAAAACCAGTAGCAAAGTTGGAAGGACAGGTA GAGCACTACTACAAAGGGGTCACAGCAAAAATTGACTACAGTAAAGGAAAAAT GCTCTTGGCCACTGACAAGTGGGAGGTGGAACATGGTGTCATAACCAGGTTAG CTAAGAGATATACTGGGGTCGGGTTCAATGGTGCATACTTAGGTGACGAGCCC AATCACCGTGCTCTAGTGGAGAGGGACTGTGCAACTATAACCAAAAACACAGT ACAGTTTCTAAAAATGAAGAAGGGGTGTGCGTTCACCTATGACCTGACCATCTC CAATCTGACCAGGCTCATCGAACTAGTACACAGGAACAATCTTGAAGAGAAGG

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GTCATGGGGGTTGGGACTGAGAGATACAAGTTAGGTCCCATAGTCAATCTGCT GCTGAGAAGGTTGAAAATTCTGCTCATGACGGCCGTCGGCGTCAGCAGCTGAg acaanatgtatatattgtaaataaattaatccatgtacAATTCCGCCCCTCTCCCTCCCCCCCTAACG TTACTGGCCGAAGCCGCTTGGAATAAGGCCGGTGTGCGTTTGTCTATATGTTAT TTTCCACCATATTGCCGTCTTTTGGCAATGTGAGGGCCCGGAAACCTGGCCCTG TCTTCTTGACGAGCATTCCTAGGGGTCTTTCCCCTCTCGCCAAAGGAATGCAAG GTCTGTTGAATGTCGTGAAGGAAGCAGTTCCTCTGGAAGCTTCTTGAAGACAAA CAACGTCTGTAGCGACCCTTTGCAGGCAGCGGAACCCCCCACCTGGCGACAGG TGCCTCTGCGGCCAAAAGCCACGTGTATAAGATACACCTGCAAAGGCGGCACA ACCCCAGTGCCACGTTGTGAGTTGGATAGTTGTGGAAAGAGTCAAATGGCTCT CCTCAAGCGTATTCAACAAGGGCTGAAGGATGCCCAGAAGGTACCCCATTGT ATGGGATCTGATCTGGGGCCTCGGTGCACATGCTTTACATGTGTTTAGTCGAG GTTAAAAAACGTCTAGGCCCCCGAACCACGGGGACGTGGTTTTCCTTTGAAA AACACGATGATAAGCTTGCCACAACcatgaccgagtacaagcccacggtgcgcctcgccacccgcgacga cgtcccccgggccgtacgcaccctcgccgccgttcgccgactaccccgccacacgcgccacaccgtcgacccggaccgccacatc gagogggtcaccgagotgcaagaactcttcctcacgcgcgtcgggctcgacatcggcaaggtgtgggtcgcgacgacggcgcc cggttcccggctggccgcgcagcaacagatggaaggctcctggcgccgcaccggcccaaggagcccgcgtggttcctggccac cgtcggcgtctcgcccgaccaccagggcaagggtctgggcagcgccgtcgtgctccccggagtggaggcggccgagcgcgccg gggtgcccgccttcctggagacctccgcgccccgcaacctccccttctacgagcggctcggcttcaccgtcaccgccgacgtcgagt AGCATTATGAGCAACTGCAGCTAAGAACAGAGACAAACCCGGTCATGGGGGTT GGGACTGAGAGATACAAGTTAGGTCCCATAGTCAATCTGCTGCTGAGAAGGTT GAAAATTCTGCTCATGACGGCCGTCGGCGTCAGCAGCTGAgacaaaatgtatattgtaaata aattaatccatgtacatagtgtatataaatatagttgggaccgtccacctcaagaagacgacacgcccaacacgcacagctaaacagtag tcaagattatctacctcaagataacactacatttaatgcacacagcactttagctgtatgaggatacgcccgacgtctatagttggactagg gaagacctctaacagccccc

Bicistronic HCV/BVDV chimera

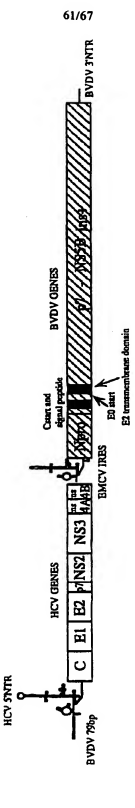


FIGURE 25

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CGGAGCGGTCTACGCCTTCTACGGGATGTGGCCTCTCCTCCTGCTCCTGCTGG CGTTGCCTCAGCGGGCATACGCACTGGACACGGAGGTGGCCGCGTCGTGG CGGCGTTGTTCTTGTCGGGTTAATGGCGCTGACTCTGTCGCCATATTACAAGCG CTACATCAGCTGGTGCATGTGGTGGCTTCAGTATTTTCTGACCAGAGTAGAAGC GTCATCTTACTCATGTGTTGTACACCCGACTCTGGTATTTGACATCACCAAAC TACTCCTGGCCATCTTCGGACCCCTTTGGATTCTTCAAGCCAGTTTGCTTAAAGT CCCTACTTCGTGCGCGTTCAAGGCCTTCTCCGGATCTGCGCGCTAGCGCGGA AGATAGCCGGAGGTCATTACGTGCAAATGGCCATCATCAAGTTAGGGGCGCTT ACTGGCACCTATGTGTATAACCATCTCACCCCTCTTCGAGACTGGGCGCACAAC GGCCTGCGAGATCTGGCCGTGGCTGTGGAACCAGTCGTCTTCTCCCGAATGGA GACCAAGCTCATCACGTGGGGGGCAGATACCGCCGCGTGCGGTGACATCATC AACGCTTGCCCGTCTCTGCCCGTAGGGGCCAGGAGATACTGCTTGGGCCAGC CGACGGAATGGTCTCCAAGGGGTGGAGGTTGCTGGCGCCCATCACGGCGTAC GCCCAGCAGACGAGAGGCCTCCTAGGGTGTATAATCACCAGCCTGACTGGCCG GGACAAAAACCAAGTGGAGGTGAGGTCCAGATCGTGTCAACTGCTACCCAAA CCTTCCTGGCAACGTGCATCAATGGGGTATGCTGGACTGTCTACCACGGGGCC GGAACGAGGACCATCGCATCACCCAAGGGTCCTGTCATCCAGATGTATACCAA TGTGGACCAAGACCTTGTGGGCTGGCCCGCTCCTCAAGGTTCCCGCTCATTGA CACCTGCACCTGCGGCTCCTCGGACCTTTACCTGGTCACGAGGCACGCCGAT GTCATTCCCGTGCGCCGGCGAGGTGATAGCAGGGGTAGCCTGCTTTCGCCCCG GCCCATTTCCTACTTGAAAGGCTCCTCGGGGGGTCCGCTGTTGTGCCCCGCGG GACACGCCGTGGGCCTATTCAGGGCCGCGGTGTGCACCCGTGGAGTGGCTAA GGCGGTGGACTTTATCCCTGTGGAGAACCTAGAGACAACCATGAGATCCCCGG TGTTCACGGACAACTCCTCTCCACCAGCAGTGCCCCAGAGCTTCCAGGTGGCC CACCTGCATGCTCCCACCGGCAGCGGTAAGAGCACCAAGGTCCCGGCTGCGTA CGCAGCCCAGGGCTACAAGGTGTTGGTGCTCAACCCCTCTGTTGCTGCAACGC TGGGCTTTGGTGCTTACATGTCCAAGGCCCATGGGGTTGATCCTAATATCAGGA CCGGGGTGAGAACAATTACCACTGGCAGCCCCATCACGTACTCCACCTACGGC **AAGTTCCTTGCCGACGGCGGGTGCTCAGGAGGTGCTTATGACATAATAATTTGT** GACGAGTGCCACTCCACGGATGCCACATCCATCTTGGGCATCGGCACTGTCCT TGACCAAGCAGAGACTGCGGGGGGGGAGACTGGTTGTGCTCGCCACTGCTACC CCTCCGGGCTCCGTCACTGTCCCATCCTAACATCGAGGAGGTTGCTCTGTCC ACCACCGGAGAGATCCCCTTTTACGGCAAGGCTATCCCCCTCGAGGTGATCAA GGGGGAAGACATCTCATCTTCTGCCACTCAAAGAAGAAGTGCGACGAGCTCG CCGCGAAGCTGGTCGCATTGGGCATCAATGCCGTGGCCTACTACCGCGGTCTT GACGTGTCTGTCATCCCGACCAGCGGCGATGTTGTCGTCGTGTCGACCGATGC TCTCATGACTGGCTTTACCGGCGACTTCGACTCTGTGATAGACTGCAACACGTG TGTCACTCAGACAGTCGATTTCAGCCTTGACCCTACCTTTACCATTGAGACAAC CACGCTCCCCAGGATGCTGTCTCCAGGACTCAACGCCGGGGCAGGACTGGC AGGGGGAAGCCAGGCATCTACAGATTTGTGGCACCGGGGGAGCGCCCCTCCG GCATGTTCGACTCGTCCGTCCTCTGTGAGTGCTATGACGCGGGCTGTGCTTGG TATGAGCTCACGCCCGCCGAGACTACAGTTAGGCTACGAGCGTACATGAACAC CCCGGGGCTTCCCGTGTGCCAGGACCATCTTGAATTTTGGGAGGGCGTCTTTA CGGGCCTCACTCATATAGATGCCCACTTTCTATCCCAGACAAAGCAGAGTGGG GAGAACTTTCCTTACCTGGTAGCGTACCAAGCCACCGTGTGCGCTAGGGCTCA AGCCCTCCCCATCGTGGGACCAGATGTGGAAGTGTTTGATCCGCCTTAAAC CCACCCTCCATGGGCCAACACCCCTGCTATACAGACTGGGCGCTGTTCAGAAT GACCTGGAGGTCGTCACGAGCACCTGGGTGCTCGTTGGCGGCGTCCTGGCTG CTCTGGCCGCGTATTGCCTGTCAACAGGCTGCGTGGTCATAGTGGGCAGGATT GTCTTGTCCGGGAAGCCGGCAATTATACCTGACAGGGAGGTTCTCTACCAGGA GTTCGATGAGATGGAAGAGTGCTCTCAGCACTTACCGTACATCGAGCAAGGGA TGATGCTCGCTGAGCAGTTCAAGCAGAAGGCCCTCGGCCTCCTGCAGACCGCG

TCCCGCCAAGCAGAGGTTATCACCCCTGCTGTCCAGACCAACTGGCAGAAACT CGAGGTCTTCTGGGCGAAGCACATGTGGAATTTCATCAGTGGGATACAATACTT GGCGGGCCTGTCAACGCTGCCTGGTAACCCCGCCATTGCTTCATTGATGGCTTT TACAGCTGCCGTCACCAGCCCACTAACCACTGGCCAAACCCTCCTCTTCAACAT ATTGGGGGGGTGGGTGCCCAGCTCGCCGCCCCGGTGCCGCTACCGCC TTTGTGGGCGCTGGCTTAGCTGGCGCCCCATCGGCAGCGTTGGACTGGGGA AGGTCCTCGTGGACATTCTTGCAGGGTATGGCGCGGGCGTGGCGGAGCTCT TGTAGCCTTCAAGATCATGAGCGGTGAGGTCCCCTCCACGGAGGACCTGGTCA ATCTGCTGCCCGCCATCCTCTCGCCTGGAGCCCTTGTAGTCGGTGTGGTCTGC GCAGCAATACTGCGCCGGCACGTTGGCCCGGGCGAGGGGGCAGTGCAATGGA TGAACCGGCTAATAGCCTTCGCCTCCCGGGGGAACCATGTTTCCCCCACGCAC TACGTGCCGGAGAGCGATGCAGCCGCCCGCGTCACTGCCATACTCAGCAGCCT CACTGTAACCCAGCTCCTGATcgCTAGaccatggggtaccgagCGTTACTGGCCGAAGCCGCTTGGAATAAGGCCGGTGTGCGTTTGTCTATATGTTATTTTCCACCATATTGCC GTCTTTTGGCAATGTGAGGGCCCGGAAACCTGGCCCTGTCTTCTTGACGAGCA TTCCTAGGGGTCTTTCCCCTCTCGCCAAAGGAATGCAAGGTCTGTTGAATGTCG TGAAGGAAGCAGTTCCTCTGGAAGCTTCTTGAAGACAACAACGTCTGTAGCG ACCTTTGCAGGCAGCGGAACCCCCCACCTGGCGACAGGTGCCTCTGCGGCCA AAAGCCACGTGTATAAGATACACCTGCAAAGGCGGCACAACCCCAGTGCCACG TTGTGAGTTGGATAGTTGTGGAAAGAGTCAAATGGCTCTCCTCAAGCGTATTCA ACAAGGGGCTGAAGGATGCCCAGAAGGTACCCCATTGTATGGGATCTGATCTG GGGCCTCGGTGCACATGCTTTACATGTGTTTAGTCGAGGTTAAAAAACGTCTAG GCCCCCGAACCACGGGACGTGGTTTTCCTTTGAAAAACACGATGATAATAT GGTGGAGGAACCTGTTTATGATCAGGCAGGTGATCCCTTATTTGGTGAAAGGG GAGCAGTCCACCTCAATCGACGCTAAAGCTCCCACACAAGAGAGGGGGAACGC GATGTTCCAACCAACTTGGCATCCTTACCAAAAAGAGGTGACTGCAGGTCGGG TAATAGCAGAGGACCTGTGAGCGGGATCTACCTGAAGCCAGGGCCACTATTTT ACCAGGACTATAAAGGTCCCGTCTATCACAGGGCCCCGCTGGAGCTCTTTGAG GAGGGATCCATGTGTGAAACGACTAAACGGATAGGGAGAGTAACTGGAAGTG GTGCCACGAGAAGTTACCAAAGGGTGTTCAGGTGGGTCCATAATAGGCTTGAC TGCCCTCTATGGGTCACAAGTTGCTCAGACACGAAAGAAGAAGAGGGAGCAACA8Ag cttGCATTGTTGGCGTGGGCAATAATAGCTATAGTTTTGTTTCAAGTTACAATGGG AGAAAACATAACACAGTGGAACctgcagTGGTTTGACCTGGAGGTGACTGACCAT CACCGGGATTACTTCGCTGAGTCCATATTAGTGGTGGTAGTAGCCCTCTTGGGT GGCAGATATGTACTTTGGTTACTGGTTACATACATGGTCTTATCAGAACAGAAG GCCTTAGGGATTCAGTATGGATCAGGGGAAGTGGTGATGATGGGCAACTTGCT **AACCCATAACAATATTGAAGTGGTGACATACTTCTTGCTGCTGTACCTACTGCT** GAGGGAGGAGCGTAAAGAAGTGGGTCTTACTCTTATACCACATCTTAGTGG TACACCCAATCAAATCTGTAATTGTGATCCTACTGATGATTGGGGATGTGGTAA AGGCCGATTCAGGGGGCCAAGAGTACTTGGGGAAAATAGACCTCTGTTTTACA ACAGTAGTACTAATCGTCATAGGTTTAATCATAGCTAGGCGTGACCCAACTATA GTGCCACTGGTAACAATAATGGCAGCACTGAGGGTCACTGAACTGACCCACCA GCCTGGAGTTGACATCGCTGTGGCGGTCATGACTATAACCCTACTGATGGTTA GCTATGTGACAGATTATTTTAGATATAAAAAATGGTTACAGTGCATTCTCAGCCT GGTATCTGGGGTGTTCTTGATAAGAAGCCTAATATACCTAGGTAGAATCGAGAT GCCAGAGGTAACTATCCCAAACTGGAGACCACTAACTTTAATACTATTATATTTG GCAATGTGTGCCTATCTTATTGCTGGTCACAACCTTGTGGGCCGACTTCTTAAC CCTAATACTGATCCTGCCTACCTATGAATTGGTTAAATTATACTATCTGAAAACT GTTAGGACTGATATAGAAAGAAGTTGGCTAGGGGGGATAGACTATACAAGAGT TGACTCCATCTACGACGTTGATGAGAGTGGAGAGGGCGTATATCTTTTTCCATC AAGGCAGAAAGCACAGGGGAATTTTTCTATACTCTTGCCCCTTATCAAAGCAAC

ACTGATAAGTTGCGTCAGCAGTAAATGGCAGCTAATATACATGAGTTACTTAAC TTTGGACTTTATGTACTACATGCACAGGAAAGTTATAGAAGAGATCTCAGGAGG TACCAACATAATATCCAGGITAGTGGCAGCACTCATAGAGCTGAACTGGTCCAT GGAAGAAGAGGAGCAAAGGCTTAAAGAAGTTTTATCTATTGTCTGGAAGGT TGAGAAACCTAATAATAAAACATAAGGTAAGGAATGAGACCGTGGCTTCTTGGT ACGGGGAGGAGGAAGTCTACGGTATGCCAAAGATCATGACTATAATCAAGGCC AGTACACTGAGTAAGAGCAGGCACTGCATAATATGCACTGTATGTGAGGGCCG AGAGTGGAAAGGTGCCCCCCCAAAATGTGGACGCCATGGGAAGCCGATA ACGTGTGGGATGTCGCTAGCAGATTTTGAAGAAGACACTATAAAAGAATCTTT ATAAGGGAAGGCAACTTTGAGGGTATGTGCAGCCGATGCCAGGGAAAGCATA GGAGGTTTGAAATGGACCGGGAACCTAAGAGTGCCAGATACTGTGCTGAGTGT **AATAGGCTGCATCCTGCTGAGGAAGGTGACTTTTGGGCAGAGTCGAGCATGTT** GGGCCTCAAAATCACCTACTTTGCGCTGATGGATGGAAAGGTGTATGATATCAC AGAGTGGGCTGGATGCCAGCGTGTGGGAATCTCCCCAGATACCCACAGAGTCC TTGTACAATATACCGCTAGGGGGCAACTATTTCTGAGAAACTTGCCCGTACTGG CAACTAAAGTAAAAATGCTCATGGTAGGCAACCTTGGAGAAGAAATTGGTAATC TGGAACATCTTGGGTGGATCCTAAGGGGGCCTGCCGTGTGTAAGAAGATCACA GAGCACGAAAAATGCCACATTAATATACTGGATAAACTAACCGCATTTTTCGGG ATCATGCCAAGGGGGACTACACCCAGAGCCCCGGTGAGGTTCCCTACGAGCTT GGGATAAGTTCAGTCGACCATGTAACCGCCGGAAAAGATCTACTGGTCTGTGA CAGCATGGGACGAACTAGAGTGGTTTGCCAAAGCAACAACAGGTTGACCGATG AGACAGAGTATGGCGTCAAGACTGACTCAGGGTGCCCAGACGGTGCCAGATG TTATGTGTTAAATCCAGAGGCCGTTAACATATCAGGATCCAAAGGGGCAGTCGT TCACCTCCAAAAGACAGGTGGAGAATTCACGTGTGTCACCGCATCAGGCACAC CGGCTTTCTTCGACCTAAAAAACTTGAAAGGATGGTCAGGCTTGCCTATATTTG AAGCCTCCAGCGGGAGGGTGGTTGGCAGAGTCAAAGTAGGGAAGAATGAAGA GTCTAAACCTACAAAAATAATGAGTGGAATCCAGACCGTCTCAAAAAACACAGC AGACCTGACCGAGATGGTCAAGAAGATAACCAGCATGAACAGGGGAGACTTCA AGCAGATTACTTTGGCAACAGGGGCAGGCAAAACCACAGAACTCCCAAAAGCA GTTATAGAGGAGATAGGAAGACACAAGAGAGTATTAGTTCTTATACCATTAAGG GCAGCGCAGAGTCAGTCTACCAGTATATGAGATTGAAACACCCAAGCATCTC TTTTAACCTAAGGATAGGGGACATGAAAGAGGGGGGACATGGCAACCGGGATA ACCTATGCATCATACGGGTACTTCTGCCAAATGCCTCAACCAAAGCTCAGAGCT GCTATGGTAGAATACTCATACATATTCTTAGATGAATACCATTGTGCCACTCCTG AACAACTGGCAATTATCGGGAAGATCCACAGATTTTCAGAGAGTATAAGGGTT GTCGCCATGACTGCCACGCCAGCAGGGTCGGTGACCACAACAGGTCAAAAGC ACCCAATAGAGGAATTCATAGCCCCCGAGGTAATGAAAGGGGAGGATCTTGGT AGTCAGTTCCTTGATATAGCAGGGTTAAAAATACCAGTGGATGAGATGAAAGG CAATATGTTGGTTTTTGTACCAACGAGAAACATGGCAGTAGAGGTAGCAAAGA AGCTAAAAGCTAAGGGCTATAACTCTGGATACTATTACAGTGGAGAGGATCCA GCCAATCTGAGAGTTGTGACATCACAATCCCCCTATGTAATCGTGGCTACAAAT GCTATTGAATCAGGAGTGACACTACCAGATTTGGACACGGTTATAGACACGGG GTTGAAATGTGAAAAGAGGGTGAGGGTATCATCAAAGATACCCTTCATCGTAA CAGGCCTTAAGAGGATGGCCGTGACTGTGGGTGAGCAGGCGCAGCGTAGGGG CAGAGTAGGTAGAGTGAAACCCGGGAGGTATTATAGGAGCCAGGAAACAGCA ACAGGGTCAAAGGACTACCACTATGACCTCTTGCAGGCACAAAGATACGGGAT TGAGGATGGAATCAACGTGACGAAATCCTTTAGGGAGATGAATTACGATTGGA GCCTATACGAGGAGGACAGCCTACTAATAACCCAGCTGGAAATACTAAATAATC TACTCATCTCAGAAGACTTGCCAGCCGCTGTTAAGAACATAATGGCCAGGACTG ATCACCCAGAGCCAATCCAACTTGCATACAACAGCTATGAAGTCCAGGTCCCG GTCCTGTTCCCAAAAATAAGGAATGGAGAAGTCACAGACACCTACGAAAATTAC TCGTTTCTAAATGCCAGAAAGTTAGGGGAGGATGTGCCCGTGTATATCTACGCT

ACTGAAGATGAGGATCTGGCAGTTGACCTCTTAGGGCTAGACTGGCCTGATCC TGGGAACCAGCAGGTAGTGGAGACTGGTAAAGCACTGAAGCAAGTGACCGGG TTGTCCTCGGCTGAAAATGCCCTACTAGTGGCTTTATTTGGGTATGTGGGTTAC CAGGCTCTCTCAAAGAGGCATGTCCCAATGATAACAGACATATATACCATCGAG GACCAGAGACTAGAAGACACCACCCACCTCCAGTATGCACCCAACGCCATAAA AACCGATGGGACAGAGACTGAACTGAAAGAACTGGCGTCGGGTGACGTGGAA **AAAATCATGGGAGCCATTTCAGATTATGCAGCTGGGGGACTGGAGTTTGTTAA** ATCCCAAGCAGAAAAGATAAAAACAGCTCCTTTGTTTAAAGAAAACGCAGAAGC CGCAAAAGGGTATGTCCAAAAATTCATTGACTCATTAATTGAAAATAAAGAAGA **AATAATCAGATATGGTTTGTGGGGAACACACACAGCACTATACAAAAGCATAGC** TGCAAGACTGGGGCATGAAACAGCGTTTGCCACACTAGTGTTAAAGTGGCTAG CTTTTGGAGGGGAATCAGTGTCAGACCACGTCAAGCAGGCGGCAGTTGATTTA GTGGTCTATTATGTGATGAATAAGCCTTCCTTCCCAGGTGACTCCGAGACACAG CAAGAAGGGAGGCGATTCGTCGCAAGCCTGTTCATCTCCGCACTGGCAACCTA CACATACAAAACTTGGAATTACCACAATCTCTCTAAAGTGGTGGAACCAGCCCT GGCTTACCTCCCCTATGCTACCAGCGCATTAAAAATGTTCACCCCAACGCGGCT GGAGAGCGTGGTGATACTGAGCACCACGATATATAAAACATACCTCTCTATAAG GAAGGGGAAGAGTGATGGATTGCTGGGTACGGGGATAAGTGCAGCCATGGAA ATCCTGTCACAAAACCCAGTATCGGTAGGTATATCTGTGATGTTGGGGGTAGG GGCAATCGCTGCGCACAACGCTATTGAGTCCAGTGAACAGAAAAGGACCCTAC TTATGAAGGTGTTTGTAAAGAACTTCTTGGATCAGGCTGCAACAGATGAGCTGG TAAAAGAAAACCCAGAAAAAATTATAATGGCCTTATTTGAAGCAGTCCAGACAA TTGGTAACCCCCTGAGACTAATATACCACCTGTATGGGGTTTACTACAAAGGTT GGGAGGCCAAGGAACTATCTGAGAGGACAGCAGGCAGAAACTTATTCACATTG ATAATGTTTGAAGCCTTCGAGTTATTAGGGATGGACTCACAAGGGAAAATAAG GAACCTGTCCGGAAATTACATTTTGGATTTGATATACGGCCTACACAAGCAAAT CAACAGAGGCTGAAGAAAATGGTACTGGGGTGGGCCCCTGCACCCTTTAGTT **GTGACTGGACCCCTAGTGACGAGAGGATCAGATTGCCAACAGACAACTATTTG** AGGGTAGAAACCAGGTGCCCATGTGGCTATGAGATGAAAGCTTTCAAAAATGT AGGTGGCAAACTTACCAAAGTGGAGGAGAGCGGGCCTTTCCTATGTAGAAACA GACCTGGTAGGGGACCAGTCAACTACAGAGTCACCAAGTATTACGATGACAAC CTCAGAGAGATAAAACCAGTAGCAAAGTTGGAAGGACAGGTAGAGCACTACTA CAAAGGGGTCACAGCAAAAATTGACTACAGTAAAGGAAAAATGCTCTTGGCCA CTGACAAGTGGGAGGTGGAACATGGTGTCATAACCAGGTTAGCTAAGAGATAT ACTGGGGTCGGGTTCAATGGTGCATACTTAGGTGACGAGCCCAATCACCGTGC TCTAGTGGAGAGGGACTGTGCAACTATAACCAAAAACACAGTACAGTTTCTAAA AATGAAGAAGGGGTGTGCGTTCACCTATGACCTGACCATCTCCAATCTGACCA GGCTCATCGAACTAGTACACAGGAACAATCTTGAAGAGAAGGAAATACCCACC TATAAAACCAGTACTAGGAGAGAGAGTAATCCCCGACCCTGTAGTTGATATCAA TTTACAACCAGAGGTGCAAGTGGACACGTCAGAGGTTGGGATCACAATAATTG GAAGGGAAACCCTGATGACAACGGGAGTGACACCTGTCTTGGAAAAAGTAGA GCCTGACGCCAGCGACAACCAAAACTCGGTGAAGATCGGGTTGGATGAGGGT **AATTACCCAGGGCCTGGAATACAGACACATACACTAACAGAAGAAATACACAA** CAGGGATGCGAGGCCCTTCATCATGATCCTGGGCTCAAGGAATTCCATATCAA ATAGGGCAAAGACTGCTAGAAATATAAATCTGTACACAGGAAATGACCCCAGG GAAATACGAGACTTGATGGCTGCAGGGCGCATGTTAGTAGTAGCACTGAGGGA TGTCGACCCTGAGCTGTCTGAAATGGTCGATTTCAAGGGGACTTTTTTAGATAG GGAGGCCTGGAGGCTCTAAGTCTCGGGCAACCTAAACCGAAGCAGGTTACCA AGGAAGCTGTTAGGAATTTGATAGAACAGAAAAAAGATGTGGAGATCCCTAAC TGGTTTGCATCAGATGACCCAGTATTTCTGGAAGTGGCCTTAAAAAATGATAAG **TACTACTTAGTAGGAGATGTTGGAGAGGTAAAAGATCAAGCTAAAGCACTTGG** GGCCACGGATCAGACAAGAATTATAAAGGAGGTAGGCTCAAGGACGTATGCCA

CACTGTTTGAGGAATTGTTGCTACGGTGCCCACCTGCAACTAAGAGCAATAAG GGGCACATGGCATCAGCTTACCAATTGGCACAGGGTAACTGGGAGCCCCTCGG TTGCGGGGTGCACCTAGGTACAATACCAGCCAGAAGGGTGAAGATACACCCAT ATGAAGCTTACCTGAAGTTGAAAGATTTCATAGAAGAAGAAGAAGAAGAAACCT AGGGTTAAGGATACAGTAATAAGAGAGCACAACAAATGGATACTTAAAAAAAT AAGGTTTCAAGGAAACCTCAACACCAAGAAAATGCTCAACCCTGGGAAACTATC TGAACAGTTGGACAGGGAGGGGCGCAAGAGGAACATCTACAACCACCAGATT GGTACTATAATGTCAAGTGCAGGCATAAGGCTGGAGAAATTGCCAATAGTGAG. GGCCCAAACCGACACCAAAACCTTTCATGAGGCAATAAGAGATAAGATAGACA AGAGTGAAAACCGGCAAAATCCAGAATTGCACAACAAATTGTTGGAGATTTTCC ACACGATAGCCCAACCCACCCTGAAACACACCTACGGTGAGGTGACGTGGGAG CAACTTGAGGCGGGATAAATAGAAAGGGGGCAGCAGGCTTCCTGGAGAAGA AGAACATCGGAGAAGTATTGGATTCAGAAAAGCACCTGGTAGAACAATTGGTC AGGGATCTGAAGGCCGGGAGAAAGATAAAAATATTATGAAACTGCAATACCAAA **AAATGAGAAGAGATGTCAGTGATGACTGGCAGGCAGGGGACCTGGTGGTT** GAGAAGAGGCCAAGAGTTATCCAATACCCTGAAGCCAAGACAAGGCTAGCCAT CACTAAGGTCATGTATAACTGGGTGAAACAGCAGCCCGTTGTGATTCCAGGAT ATGAAGGAAAGACCCCCTTGTTCAACATCTTTGATAAAGTGAGAAAGGAATGG GACTCGTTCAATGAGCCAGTGGCCGTAAGTTTTGACACCAAAGCCTGGGACAC TCAAGTGACTAGTAAGGATCTGCAACTTATTGGAGAAATCCAGAAATATTACTA TAAGAAGGAGTGGCACAAGTTCATTGACACCATCACCGACCACATGACAGAAG TACCAGTTATAACAGCAGATGGTGAAGTATATATAAGAAATGGGCAGAGAGGGG AGCGGCCAGCCAGACACAGTGCTGGCAACAGCATGTTAAATGTCCTGACAAT GATGTACGCCTTCTGCGAAAGCACAGGGGTACCGTACAAGAGTTTCAACAGGG TGGCAAGGATCCACGTCTGTGGGGATGATGGCTTCTTAATAACTGAAAAAGGG TTAGGGCTGAAATTTGCTAACAAAGGGATGCAGATTCTTCATGAAGCAGGCAA **ACCTCAGAAGATAACGGAAGGGGAAAAGATGAAAGTTGCCTATAGATTTGAGG ATATAGAGTTCTGTTCTCATACCCCAGTCCCTGTTAGGTGGTCCGACAACACCA GTAGTCACATGGCCGGGAGAGACACCGCTGTGATACTATCAAAGATGGCAACA** AGATTGGATTCAAGTGGAGAGAGGGGTACCACAGCATATGAAAAAGCGGTAG CCTTCAGTTTCTTGCTGATGTATTCCTGGAACCCGCTTGTTAGGAGGATTTGCCT GTTGGTCCTTTCGCAACAGCCAGAGACAGACCCATCAAAACATGCCACTTATTA TTACAAAGGTGATCCAATAGGGGCCTATAAAGATGTAATAGGTCGGAATCTAA GTGAACTGAAGAGAACAGGCTTTGAGAAATTGGCAAATCTAAACCTAAGCCTG TCCACGTTGGGGATCTGGACTAAGCACAAGCAAAAGAATAATTCAGGACTG TGTTGCCATTGGGAAAGAAGAGGGCAACTGGCTAGTTAACGCCGACAGGCTGA TATCCAGCAAAACTGGCCACTTATACATACCTGATAAAGGCTTTACATTACAAG GAAAGCATTATGAGCAACTGCAGCTAAGAACAGAGACAAACCCGGTCATGGGG GTTGGGACTGAGAGATACAAGTTAGGTCCCATAGTCAATCTGCTGCTGAGAAG **GTTGAAAATTCTGCTCATGACGGCCGTCGGCGTCAGCAGCTGAgacaaaatgtatatattgt** anatanattaniccatgincatagigtatatanaatatagiiggaccgiccaccicangangacgacacgcccaacacgcacagcinnac agtagtcaagattatctacctcaagataacactacatttaatgcacacagcactttagctgtatgaggatacgcccgacgtctatagttggac tagggaagacctctaacagccccc

INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/08850

IPC(6) :	SSIFICATION OF SUBJECT MATTER A61K 39/29, 39/295; C12Q 1/70; C12N 7/01; C07H 2 424/218.1, 228.1; 435/5, 235.1; 536/23.72		·						
	o International Patent Classification (IPC) or to both n	ational classification and ire							
B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols)									
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U.S. : 4	424/218.1, 228.1; 435/5, 235.1; 536/23.72								
Documentati	ion searched other than minimum documentation to the	extent that such documents are included	in the fields searched						
Electronic d	ata base consulted during the international search (nar	ne of data base and, where practicable,	search terms used)						
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C. DOC	UMENTS CONSIDERED TO BE RELEVANT		· · · · · · · · · · · · · · · · · · ·						
Category*	Citation of document, with indication, where app	ropriate, of the relevant passages	Relevant to claim No.						
X,P	FROLOV et al. cis-acting RNA elemer bovine viral diarrhea virus-hepatitis Cychimeras. RNA. November 1998, Ventire document.	virus 5' nontranslated region	1-8, 10-21						
Y,P	MALET et al. Yellow fever 5' noncelement to improve hepatitis C virus proof translational control. Biochem. Biochem. Biochem. Biochem. Biochem. 253, No. 2, document.	duction through modification iophys. Res. Commun. 18							
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International application No.
PCT/US99/08850

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	LU et al. Poliovirus chimeras replicating under the translational control of genetic elements of hepatitis C virus reveal unusual properties of the internal ribosomal entry site of hepatitis C virus. Proc. Natl. Acad. Sci. USA. 20 February 1996, Vol. 93, No. 4, pages 1412-1417, see entire document.	1-8, 10-21
Y	VASSILEV et al. Authentic and chimeric full-length genomic cDNA clones of bovine viral diarrhea virus that yield infectious transcripts. J. Virol. January 1997, Vol. 71, No. 1, pages 471-478, see entire document.	1-8, 10-21
Y	VENUGOPAL et al. Towards a new generation of flavivirus vaccines. Vaccines. 1994, Vol. 12, No. 11, pages 966-975, see entire document.	1-8, 10-21
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/08850

Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: 9 because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically: CLAIM 9 RECITES "SEQ ID NO:X" WHICH EXPRESSION IS NOT UNDERSTOOD.
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

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